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PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

NEWS Web Page for STN Seminar Schedule - N. America NEWS NOV 21 CAS patent coverage to include exemplified prophetic substances identified in English-, French-, German-, and Japanese-language basic patents from 2004-present NEWS NOV 26 MARPAT enhanced with FSORT command NEWS NOV 26 CHEMSAFE now available on STN Easy NEWS NOV 26 Two new SET commands increase convenience of STN searching DEC 01 ChemPort single article sales feature unavailable NEWS 6 NEWS DEC 12 GBFULL now offers single source for full-text coverage of complete UK patent families DEC 17 Fifty-one pharmaceutical ingredients added to PS NEWS 8

NEWS 9 JAN 06 The retention policy for unread STNmail messages will change in 2009 for STN-Columbus and STN-Tokyo

NEWS 10 JAN 07 WPIDS, WPINDEX, and WPIX enhanced Japanese Patent Classification Data

NEWS 11 FEB 02 Simultaneous left and right truncation (SLART) added for CERAB, COMPUAB, ELCOM, and SOLIDSTATEM

NEWS 12 FEB 02 GENBANK enhanced with SET PLURALS and SET SPELLING

NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3, AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

NEWS HOURS STN Operating Hours Plus Help Desk Availability

NEWS LOGIN Welcome Banner and News Items

NEWS IPC8 For general information regarding STN implementation of IPC 8

Enter NEWS followed by the item number or name to see news on that specific topic.

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FILE 'HOME' ENTERED AT 08:03:02 ON 02 FEB 2009

=> FIL REG

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.22 0.22

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 08:03:21 ON 02 FEB 2009
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 30 JAN 2009 HIGHEST RN 1098270-10-0 DICTIONARY FILE UPDATES: 30 JAN 2009 HIGHEST RN 1098270-10-0

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH July 5, 2008.

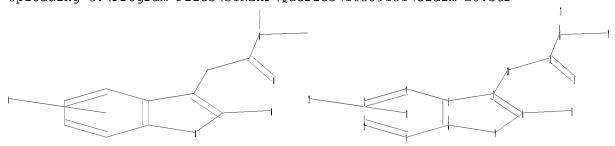
Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=>

Uploading C:\Program Files\STNEXP\Queries\10539151\claim 20.str



chain nodes :

10 11 12 14 15

ring nodes :

1 2 3 4 5 6 7 8 9

ring/chain nodes :

16 17 18

chain bonds :

7-10 8-11 10-14 14-15 14-16

ring/chain bonds :

16-17 16-18

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9

exact/norm bonds :

5-7 6-9 7-8 8-9 14-15 14-16 16-17 16-18

exact bonds: 7-10 8-11 10-14 normalized bonds:

1-2 1-6 2-3 3-4 4-5 5-6

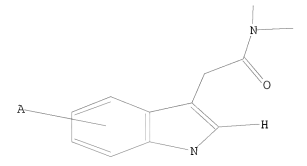
Match level:

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:Atom 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS

L1 STRUCTURE UPLOADED

=> D

L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> S L1

SAMPLE SEARCH INITIATED 08:03:47 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 2843 TO ITERATE

70.3% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 53662 TO 60058 PROJECTED ANSWERS: 869 TO 1859

L2 48 SEA SSS SAM L1

=> D SCAN

48 ANSWERS

48 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN INDEX NAME NOT YET ASSIGNED C29 H33 N3 06

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

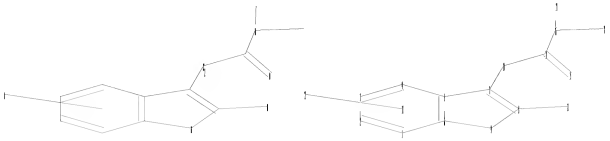
48 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN 1H-Indole-7-carboximidamide, N-hydroxy-3-[2-[4-(1-isoquinoliny1)-1-piperaxiny1]-2-oxoacety1]-4-methoxy- C25 H24 N6 O4

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=>

Uploading C:\Program Files\STNEXP\Queries\10539151\claim 20_2.str



chain nodes : 10 11 12 14 15 ring nodes : 1 2 3 4 5 6 7 8 9 ring/chain nodes : 16 17 18 chain bonds : 7-10 8-11 10-14 14-15 14-16 ring/chain bonds : 16-17 16-18 ring bonds : 1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9 exact/norm bonds : 5-7 6-9 7-8 8-9 14-15 14-16 16-17 16-18 exact bonds : 7-10 8-11 10-14

Match level :

normalized bonds :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:Atom 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS

L3 STRUCTURE UPLOADED

1-2 1-6 2-3 3-4 4-5 5-6

=> D L3 HAS NO ANSWERS L3 STR

1 ANSWERS

Structure attributes must be viewed using STN Express query preparation.

=> S L3

SAMPLE SEARCH INITIATED 08:05:03 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 2843 TO ITERATE

70.3% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 53662 TO 60058 PROJECTED ANSWERS: 1 TO 99

L4 1 SEA SSS SAM L3

=> D SCAN

L4 1 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN

IN 1H-Indole-3-acetamide, 5-bromo-N-cyclopentyl-N-[2-[5-methyl-3-(4pyridinyl)-1H-1,2,4-triazol-1-yl]ethyl]
MF C25 H27 Br N6 0

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=> S L3 FULL

FULL SEARCH INITIATED 08:05:12 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 55859 TO ITERATE

100.0% PROCESSED 55859 ITERATIONS

152 ANSWERS

SEARCH TIME: 00.00.01

L5 152 SEA SSS FUL L3

=> D SCAN

L5 152 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN
IN 1H-Indole-5-carbonitrile, 3-[2-oxo-2-(1-piperaziny1)ethy1]-, hydrochloride
(1:1)
MF C15 H16 N4 O . C1 H

● HCl

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

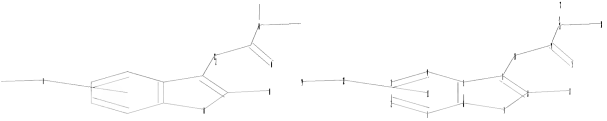
152 ANSWERS REGISTRY COPYRIGHT 2009 ACS on STN Ethanone, 2-[5-(phenylmethoxy)-1H-indol-3-y1]-1-(1-piperidinyl)-C22 H24 N2 O2

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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chain nodes :

10 11 12 14 15 19

ring nodes :

1 2 3 4 5 6 7 8 9

ring/chain nodes :

16 17 18

chain bonds :

7-10 8-11 10-14 12-19 14-15 14-16

ring/chain bonds :

16-17 16-18

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9

exact/norm bonds :

5-7 6-9 7-8 8-9 12-19 14-15 14-16 16-17 16-18

exact bonds : 7-10 8-11 10-14

normalized bonds : 1-2 1-6 2-3 3-4 4-5 5-6

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:Atom 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS

L6 STRUCTURE UPLOADED

=> D

L6 HAS NO ANSWERS

L6 STR

Structure attributes must be viewed using STN Express query preparation.

=> S L6 FULL SUB=L5
FULL SUBSET SEARCH INITIATED 08:05:52 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 152 TO ITERATE

100.0% PROCESSED 152 ITERATIONS 68 ANSWERS

SEARCH TIME: 00.00.01

L7 68 SEA SUB=L5 SSS FUL L6

=> S L6 NOT L7

L7 MAY NOT BE USED HERE

The L-number entered was not created by a STRUCTURE or SCREEN command.

=> D HIS

(FILE 'HOME' ENTERED AT 08:03:02 ON 02 FEB 2009)

FILE 'REGISTRY' ENTERED AT 08:03:21 ON 02 FEB 2009 L1 STRUCTURE UPLOADED L2 48 S L1 L3 STRUCTURE UPLOADED L41 S L3 L5 152 S L3 FULL STRUCTURE UPLOADED L6 68 S L6 FULL SUB=L5 L7 => S L5 NOT L7 84 L5 NOT L7 L8

> EII CADIIIC

=> FIL CAPLUS
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 232.76 232.98

FULL ESTIMATED COST

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FILE COVERS 1907 - 2 Feb 2009 VOL 150 ISS 6 FILE LAST UPDATED: 30 Jan 2009 (20090130/ED)

Caplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> S L8 L9 44 L8

=> D IBIB 1-5

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L9 ANSWER 2 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:581016 CAPLUS
                                                  44 CAPLUS COPYRIGHT 2009 ACS on STN 2008:1006368 CAPLUS
               ANSWER 1 C
  ACCESSION NUMBER
    DOCUMENT NUMBER:
                                                                       149:307661
                                                                                                                                                                                                                                                        DOCUMENT NUMBER:
                                                                                                                                                                                                                                                                                                                            149:104570
                                                                            ovel indole derivatives as inhibitors hepatitis C
                                                                                                                                                                                                                                                                                                                             2-Aminomethyl piperidines as novel urotensin-II
  TITLE:
                                                                                                                                                                                                                                                        TITLE:
                                                                                                                                                                                                                                                                                                                           2-Aminomethyl piperidines as novel urotensin-ii receptor antaqonists Jin, Jian; Wang, Yonghui, Wang, Feng; Shi, Dongchuan; Erhard, Karl F.; Wu, Zining, Guida, Brian F.; Lawrence, Sarah K.; Behm, David J.; Disa, Jyeti; Vaidya, Kalindi S.; Evans, Christopher; McMillan, Lynette J.; Rivezo, Ralph A.; Neeb, Michael J.; Douglas, Stephen A. GlaxoSmithKline, Cardiovascular and Urogenital Center of Excellence for Drug Discovery, King of Prussia,
                                                                        virus replication and their preparation and use in
                                                                  Andrews, Steven W.; Misialek, Shawn Maurice;
Rajaqualan, P. T. Ravi; Fryer, Andrew M.;
Madduru Machender R.; Zhang, Gan; Kossen, Karl;
Serebryan, Vladimir
Intermune, Inc., USA
PCT Int. Appl., 397pp.
CODEN; FIXXD.
Patent
                                                                                                                                                                                                                                                        AUTHOR(S):
   INVENTOR(S):
  Dimitrova
                                                                                                                                                                                                                                                       CORPORATE SOURCE:
                                                                                                                                                                                                                                                                                                                           19406, USA
Bioorganic & Medicinal Chemistry Letters (2008),
18(9), 2860-2864
CODEN: BMCLE8; ISSN: 0960-894X
Elsevier Ltd.
Journal
English
CASREACT 149:104570
26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR
  PATENT ASSIGNEE(S);
SOURCE:
                                                                                                                                                                                                                                                        SOURCE:
                                                                                                                                                                                                                                                       PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
REFERENCE COUNT:
  DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
               PATENT NO.
                                                                                         DATE
PATENT NO. KIND DATE ANDLICATION NO.

WO 2008100867 A2 20080821 WO X08-US53617

W: AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH,
CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ,
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ME, MS, MK, MN, MM, MM, MY, MZ, NA, NS, NI, NO,
PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, M,
TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZA,
RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR,
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TG, BM, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ,
AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EE,
PRIORITY APPLN. INFO:
                                                                                                                                                                                                                                                                                                                                            RECORD. ALL CITATIONS AVAILABLE IN THE RE
                                                                                                                                                 SE, SI, SK,
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                                                                                                                                                                                   NE, SN, TD,
UG, ZM, ZW,
                                                                                                                                                                                            20070212
  OTHER SOURCE(S):
                                                                    MARPAT 149:307661
  L9 ANSWER 3 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2008:505080 CAPLUS
                                                                                                                                                                                                                                                                             SWER 4 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
                                                                     LUS COPYRIGHT 2009 ACS on STN 2008;505080 CAPLUS 148:495.796 Preparation of deuterated aminoethylindolylmethylsulfonamides as serotonin 5-HTIB and/or 5-HTID receptor modulators. Gant, Thomas G., Sarshar, Sepehr Auspex Pharmaceuticals, Inc., USA PCT Int. Appl., 117pp. CODEN: PIXXD2 Patent
                                                                                                                                                                                                                                                                                                                            US COPYRIGHT 2009 ACS on STN
2008:501180 CAPLUS
148:495981
148:495981
Freparation of piperazine-substituted benzothiophenes
for treatment of mental disorders
Yamashita, Hiroshi, Matsubara, Jun; Oshima, Kunio;
Kuroda, Hideaki; Shimizu, Satoshi; Tanaka,
                                                                                                                                                                                                                                                        ACCESSI
     OCUMENT NUMBER:
                                                                                                                                                                                                                                                        INVENTOR (S)
                                                                                                                                                                                                                                                                                                                            Taira, Shinichi; Kondo, Kazumi; Takahashi, Haruka; Fukushima, Tae; Sakurai, Yohji
Otsuka Pharmaceutical Co., Ltd., Japan
PCT Int. Appl., 312pp.
CODEN: PIXXD2
Patent
English
   PATENT ASSIGNEE(S):
SOURCE:
                                                                      Patent
English
  DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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SOURCE:
         PATENT NO. KIND DATE

WO 2008049116 A2 20080424 WO 2007-US81977
WO 2008049116 A3 20080605
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BM
CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, E
GB, GD, GZ, GH, CM, GT, HN, HR, HU, ID, IL, IN, I
KM, NN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, L
MG, MK, MM, MM, MM, MM, MM, MZ, MZ, NA, NG, NI, NO, NZ, C
FT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, S
TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
RN: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, (
IS, IT, LT, LU, LV, MC, MT, ML, PL, PT, RO, SE,
GH, GM, KE, LS, MM, MZ, NM, SN, SL, SZ, TZ, UG,
BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, CA
US 20080103189 A1 20080501 US 2007-075570
US 2006-053243P
                                                                                                                                                                                                                                                        DOCUMENT TYPE:
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FAMILY ACC. NUM. CO
PATENT INFORMATION:
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                                                                                                                                                                                                                                                                                                                                                                                APPLICATION NO.
                                                                                                                                                                                                                                                                                                                                               DATE
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FI, FR, GB, GR, HU, IE,
RO, SE, SI, SK, TR, BF,
MR, NE, SN, TD, TG, BW,
TZ, UG, ZM, ZW, AM, AZ,
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  PRIORITY APPLN. INFO.:
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JP 2008115175
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                                                                    CASREACT 148:495786: MARPAT 148:495786
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  OTHER SOURCE(S):
                                                                                                                                                                                                                                                                                                                                                                                JP 200
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                                                                                                                                                                                                                                                                                                                                                                                                                                     A 20061013
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                                                                                                                                                                                                                                                        REFERENCE COUNT:
                                                                                                                                                                                                                                                                                                                                            THERE ARE 4 CITED REFEREN S AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
                                                                                                                                                                                                                                                        FORMAT
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L9 ANSWER 7 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2007:730896 CAPLUS
                    ANSWER 6 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 2007:1145534 CAPLUS
                                                                          147:448797
Preparation of aminopyrrolidine derivatives as MC4 receptor antagonists for treatment of depression, anxiety disorder, etc.
Okubo, Taketoshi; Kumagai, Toshihito; Ishii, Takaaki; Nakamura, Toshio; Abe, Kumi; Amada, Yuri; Ishizaka, Tomoko; Sun, Xiang-Min; Sekiguchi, Yoshinori; Sasako, Shigetada; Shimizu, Takanori; Nagatsuka, Takayuki Talsho Pharmaceutical Co., Ltd., Japan FCT Int. Appl., 230pp.
CODEN: PIXXD2
Patent
Japanese
                   MENT NUMBER:
                                                                                                                                                                                                                                                                 DOCUMENT NUMBER:
                                                                                                                                                                                                                                                                                                                                       147:143468
                                                                                                                                                                                                                                                                 TITLE:
                                                                                                                                                                                                                                                                                                                                       Heterocyclic derivatives as modulators of ion
                                                                                                                                                                                                                                                                 channels
                                                                                                                                                                                                                                                                                                                                      and their preparation, pharmaceutical compositions
      INVENTOR (S
                                                                                                                                                                                                                                                                and
                                                                                                                                                                                                                                                                                                                                    use in the treatment of diseases
Wilson, Dean; Fanning, Lev T. D.; Sheth, Urvi;
Martinborough, Esther; Termin, Andreas; Neubert,
Timothy; Zimmermann, Nicole; Knoll, Tara; Whitney
Tara; Kawatkar, Aarti; Lehsten, Danielle; Stamos,
Dean; Zhou, Jinglan; Arumugam, Vljayalaksmí;
Gutierrez, Corey
Vertex Pharmaceuticals Incorporated, USA
PCT Int. Appl., 369pp.
CODEN; PIXXD2
Fatent
English
1
                                                                                                                                                                                                                                                                INVENTOR (S) +
      PATENT ASSIGNEE(
      DOCUMENT TYPE:
      DOCUMENT TIPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
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LANGUAGE;
FAMILY ACC. NUM. COUNT;
PATENT INFORMATION:
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PATENT NO. KIND DATE APPLICAGE.

WO 2007075895 A2 20070705 WO 2006-US48802 20061221

WO 2007075895 A3 20071129

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CM, CO, CR, CU, CZ, DEP, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JF, KZ, KG, KM, KN, KF, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MM, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, CM, FG, PH, FL, PT, RO, RS, RU, SC, SD, SE, SC, SK, SL, SL, SM, SV, Y, IJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW

RN: AT, BE, BG, CH, CY, CZ, DE, DK, ER, ES, FI, FR, GB, GR, HU, IE, LS, TT, LT, LU, LV, MG, NL, PL, FT, RO, SE, SI, SK, TR, BR, BJ, CF, CG, CI, CM, GA, GN, CQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, FY, KG, GR, ST, ND, RT, TJ, TN, AR, EA, EF, CA

AU 2006331608 A1 20070705 CA 2006-2633653 20061221

ED 1963281 A2 20080930 EF 2006-245551 20061221

ED 1963281 A2 CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, CR, EF, EF, FI, FR, GB, GR, HU, II, MM, MZ, ER, EF, EF, FI, FR, GB, GR, HU, II, MM, EF, FI, FR, GB, GR, HU, III, MM, EF, FI, FR, GB, GR, HU, III, MM, EF, FI, FR, GB, GR, HU, III, MM, EF, FI, FR, GB, GR, 
                                                                                                                                WO 2007-JP57054
                                                                                                                                                                                                20070330
      OTHER SOURCE(S):
                                                                           MARPAT 147:448797

13 THERE ARE 13 CITED REFERENCES AVAILABLE
                                                                                                                                                                                                                                                                              EP 1963281 A2 20080903 EP 2006-845951 20061221
R: AT, BE, BE, CH, CY, CZ, DE, DK, EE, ES, FI, FE, GB, CR, HU, IE,
IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL,
BA, HR, MK, RS
MX 2008008204 A 20080715 MX 2008-8204 20080620
      REFERENCE COUNT:
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IN 2008-KN2697
KR 2008-717835
NO 2008-3220
US 2005-752926P
      FORMAT
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KR 2008081178
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     L9 ANSWER 7 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
US 2006-839444P P 20060823
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ACCESSION NUMBER:
                                                                                                                                                                                                                                                                                                       OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
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144:331420
                                                                                                                                                                                                                                                                  DOCUMENT NUMBER
                                                                                                                                                                                                                                                                                                                                       144:331420
Preparation of bicyclic anilide spirolactam cgrp receptor antagonists
Bell, Ian M.; Theberge, Cory R.; Stump, Craig A.; Zhang, Xufang; Gallicchio, Steven N.; Zartman, C. Blair.
                                                                                                                                WO 2006-US48802
                                                                                                                                                                                W 20061221
      OTHER SOURCE(S):
                                                                        MARPAT 147:143468
                                                                                                                                                                                                                                                                 INVENTOR(S):
                                                                                                                                                                                                                                                                 PATENT ASSIGNEE(S):
SOURCE:
                                                                                                                                                                                                                                                                                                                                       Merck & Co., Inc., USA
FT Int. Appl., 116 pp.
CODEN: PIXXD2
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Patent
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LANGUAGE:
FAMILY ACC. NUM. CO
PATENT INFORMATION:
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EP 1797073
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IN 2007DN01493
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IN 2007-DN1493
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                                                                                                                                                                                                                                                                                                                                                                                                                                                W 20050909
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ANSWER 9 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SION NUMBER: 2004:902086 CAPLUS
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         MENT NUMBER:
                                                      141:388753
                                                      141:388753
Heterocyclic compound modulators of Tie-2 and other kinases, and therapeutic use
Chen, Jeff; Dalrymple, Lisa; Epshteyn, Sergery;
Forsyth, Timothy; Buynh, Tai; Leahy, James; Mann,
Grace; Mann, Larry W.; Ridgway, Brian; Sangalang,
                                                                                                                                                                                                                                                           Preparation of N-heterocyclyl-substituted
amino-thiazole derivatives as protein kinase
 TITLE
                                                                                                                                                                                                     TITLE:
                                                                                                                                                                                                                                                         amino-thiazole derivatives as protein kinase inhibitors
Alegria, Larry Andrew; Chong, Wesley Kwan Mung; Chu, Shaosong, Duvadie, Rohit Kumar; Li, Lin; Romines, William Henry, III; Yang, Yi
Pfizer Inc., USA
PCT Int. Appl., 307 pp.
CODEN: FIXXD2
Fatent
English 1
INVENTOR (
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                                                      C.; Takeuchi, Craig
Exelixis, Inc., USA
CT Int. Appl., 126 pp.
COLN: PIXXD2
PateN:
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PATENT ASSIGNEE(S):
                                                                                                                                                                                                    DOCUMENT TYPE:
LANGUAGE;
FAMILY ACC. NUM, COUNT;
PATENT INFORMATION:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC, NUM. COUNT:
PATENT INFORMATION:
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          PATENT NO. KIND DATE APPLICATION NO. DI

WO 2004091480 A2 20041028 W 2004-US10626 20

W: AR, AG, AL, AM, AT, AU, AZ, BA, BB, BZ, BR, BW, BY, BZ,
GE, GH, CW, HR, HU, ID, IL, IN, IS, JP, NS, KG, KP, KR,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, WN, MX, MZ,
NO, NZ, CM, FG, PH, FL, PT, RO, RU, SC, SD, SR, SG, SK,
IJ, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN,
EY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ,
SK, TR, EF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
TD, TG
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FI, GB, GD,
KR, KZ, LC,
MZ, NA, NI,
SK, SL, SY,
ZA, ZM, ZW
ZW, AM, AZ,
E, DK, EE,
RO, SE, SI,
MR, NS, SN,
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           TD, TG
AU 2004229392
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CA 2004-2520255
EP 2004-759191
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          CA 2520255
EP 1611123
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20060104
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                          AZ 20060104 EF 2004-759191 20040408
AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, FL, SK,
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                   R:
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          JP 2006522813
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3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
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REFERENCE COUNT:
US 20060293342
PRIORITY APPLN. INFO.:
                                                                     20061228
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US 2003-461471P
                                                                                                                                          P 20030409
                                                                                               WO 2004-US10626
                                                                                                                                                                                                     FORMAT
                                                                                                                                          A 20040408
                                                      MARPAT 141:388753

1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
OTHER SOURCE(S):
REFERENCE COUNT:
FORMAT
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ANSWER 11 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 2004:546477 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 141:89009 191:03003 Synthesis of tryptamine derivatives and intermediates thereof TITLE: thereof Berens, Ulrich; Dosenbach, Oliver; Sprenger, Daniel Ciha Specialty Chemicals Holding Inc., Switz. PCT Int. Appl., 84 pp. CODEN: PIXXD2 Fatent INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION: English PATENT NO. KIND DATE APPLICATION NO. DATE W0 2004056769 A2 20040708 W0 2003-EP50992
W0 2004056769 A3 20040916
W1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CO, CR, CU, CZ, DE, DX, DM, DZ, EG, EE, EG, ES, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, LR, LS, LI, LU, LV, MA, MD, MG, MG, MK, MN, MN, MX, CM, FG, FH, FL, FT, FO, EU, SC, SD, SE, SG, SK, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, RW1 EW, GH, GM, KE, LS, MN, MZ, SD, SL, SZ, TZ, UG, BY, KG, KS, MD, RU, TO, TM, AT, BE, BG, CI, CY, ES, F1, FR, GB, GR, HU, IE, IT, LU, MC, ML, PT, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, 20031212 TG A T A1 US 20060058367 IN 2005CN01638 IN 2007CN05032 PRIORITY APPLN, INFO.: 20060316 US 2005-539151 IN 2005-CN1638 20050616 20050719

IN 2005-CN1638 A3 20050719 OTHER SOURCE(S): MARPAT 141:89009

20080321

IN 2007-CN5032 EP 2002-406128

WO 2003-EP50992

A 20021220

W 20031212

ANSWER 11 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) 717139-80-59 717139-94-99 Rt. SPN (Synthetic preparation), PREP (Preparation) (preparation of tryptamine derivs. and intermediates thereof) 717139-80-5 CAPLUS 1H-Indole-3-acetamide, 5-bromo-N,N-dimethyl-1-(phenylmethyl)- (CA INDEX NAME)

717139-84-9 CAPLUS
1H-Indole-3-acetamide, 5-iodo-N,N-dimethyl-1-(phenylmethyl)- (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 11 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Indoleacetates I [R = CO2R3; R1 = (un)substituted alkyl, aryl, heterocyclyl, alkylsulfonyl, OH, SH, NO2, halogen, CN, CONH2, CONHNH2, CO2H, alkenyl, alkynyl, cycloalkyl, acyloxy, NH2, NHNH2, B(GH)2; R2 = H, (un)substituted alkyl, CO2H, arylsulfonyl, alkylsulfonyl, aryl, CONH3, silyl; R3 = (un)substituted alkyl, n = 0-4] were prepared and converted

[R = CONR4R5; R4, R5 = (un)substituted alkyl; R4R5 = (un)substituted alkylene] which were in turn converted to indoleacetamides and tryptamines. The synthesis methods and products are useful in the synthesis of pharmaceuticals. Thus, 5-bromolsatin was treated with CH2(COZH)2 and CICONMe2 to give I [R = CONNe2, R1 = 5-Br, R2 = H] which was treated with BF3.Et20 and BH3.Me2SO to give 2-(5-bromo-1H-indol-3-yl)-N,N-dimethylacetamide or with BF3.Et20 and

4
to give [2-(5-bromo-1H-indol-3-yl)ethyl]-N,N-dimethylacetamide.
717139-79-2P 717139-83-8F
RL: RCT (Reactant); SFN (Synthetic preparation); FREP (Preparation); RACT (Reactant or reagent)
(preparation of tryptamine derivs. and intermediates thereof)
717139-79-2 CAPLUS
1H-Indole-3-acetamide, 5-bromo-N,N-dimethyl- (CA INDEX NAME)

$$\begin{array}{c|c} & H & \\ & N & \\ & & \\ & CH_2-C-NMe_2 \end{array}$$

717139-83-8 CAPLUS
1H-Indole-3-acetamide, 5-iodo-N,N-dimethyl- (CA INDEX NAME)

ACCESSION NUMBER: DOCUMENT NUMBER:

ANSWER 12 OF 44 CAPLUS COPYRIGHT 2009 ACS ON STN

SSION NUMBER: 2004:525891 CAPLUS

MENT NUMBER: 141:89110

Preparation of piperazinylethylindolecarbonitriles as serotonin reuptake inhibitors and 5-HTIA/5-HTIB receptor liquands.

NTOR(S): Heinrich, Timo; Boettcher, Henning; Schiemann, Kai; Heolzemann, Guenter; van Amsterdam, Christoph; Bartoszyk, Gerd; Leibrock, Joachim; Seyfried, Christoph

NT ASSIGNEE(S): Merck Patent GmbH, Germany

GET. OCTOBEN: GNXXSX

PATENT

UNGG: GETMAN COUNT: 1

NT INFORMATION: INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

		TENT :																
		1005																
		1025															0021	
		2510																
	MO	2004																
		W:										BG,						
												EC,						
												JP,						
												MK,						
			NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ΤJ,
			TM,	TN,	TR,	TT,	TZ,	UΑ,	ŪG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW	
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	zw,	AM,	AZ,
			BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,
			ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,
			TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,
TG																		
	ΑU	2003	2981	45		A1		2004	0709		AU 2	2003-	2981	45		2	0031	127
	EP	1572	646			A1		2005	0914		EP 2	2003-	7958	48		2	0031	127
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	ĒĒ,	HU,	SK	
	BR	2003	0174	22		A		2005	1108		BR 2	2003-	1742	2		2	0031	127
	CN	1729	173			A		2006	0201		CN 2	2003-	8010	6737		2	0031	127
	JP	2006	5115	22		T		2006	0406		JP 2	2004-	5597:	27		2	0031	127
		2005															0050	614
	US	2006	0122	191		A1		2006	0608		US 2	2005-	5395	16		2	0050	617
		2005															0050	714
PRIO	RIT	APP	LN.	INFO	. :						DE 2	2002-	1025	9244		A 2	0021	217

WO 2003-EP13374

OTHER SOURCE(S): MARPAT 141:89110

$$\underset{R111}{\overset{R11}{\overbrace{\hspace{1.5cm}}}}\underset{H}{\overset{N}{\underset{\hspace{1.5cm}\times\hspace{1.5c$$

w 20031127

ANSWER 12 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Title compds. [I; R11, R111 = H, cyano, halo, A, OA, OH, COR2, CH2R2; R2

OH, OA, NH2, NHA, NA2; h = (fluoro-substituted) alkyl optionally interrupted by O, S, CH:CH; h = (partially or completely saturated) (aubstituted) mono- or polycyclic carbo- or heterocyclyl; n = 0-4], were prepared Thus, 3-(2-chloroeth-1-yl)-lH-indole-5-carbonitrile

(substituted) mono— or polycyclic carbo— or heterocyclyl, n = 0-4], were prepared Thus, 3-(2-chloroeth-1-yl)-1H-indole-5-carbonitrile paration given), 1-(2,3-dihydrobenzo[1,4]-dioxin-5-yl)piperazine, ethyldiisopropylamine, and N-methylpyrrolidinone were heated together at 120° for 12 h to give 3-[2-[4-(2,3-dihydrobenzo[1,4]dioxin-5-yl)piperazin-1-yl]ethyl-1H-indole-5-carbonitrile. The latter showed SSRI, 5-HTIA, and 5-HTIB receptor activity at 11 nM, 17 nM, and 11 nM, resp.
714954-07-1P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation), USES (UGes)
(preparation of piperazinylethylindolecarbonitriles as serotonin

(preparation of piperazinylethylindolecarbonitriles as serotonin

take inhibitors and receptor ligands)
714954-07-1 CAPLUS
1H-Indole-5-carbonitrile, 3-[2-[4-[2-(5-fluoro-lH-indol-3-y1)acetyl]-l-piperazinyl]ethyl]-, hydrochloride (l:1) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

• HCl

ANSWER 13 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

This invention provides indole, azaindole, and related heterocyclic piperazinecarboxamides Q(C(O))m(CR8R8')n(C(O))pTC(O)A (1; variables defined below; e.g. $N-(benzoy1)-N^*-[2-(indol-2-y1)-2-oxo-1-cyanoethy]piperazine (shown as I)) having drug and blo-affecting properties, their pharmaceutical compns. and method of use. These is$

possess unique antiviral activity, whether used alone or in combination with other antivirals, antiinfectives, immunomodulators or HIV entry inhibitors. More particularly, the present invention relates to the treatment of HIV and AIDS. EC50 ranges against HIV-1 are given for about 30 of the claimed compds.; for example, N-(benzoyl)-N-(2-(6-methoxyindol-2-yl)-2-oxo-1-cyanoethyl)-3-methylpiperazine has an EC50 < lpuM. Although the methods of preparation

are

not claimed, 32 example prepns. of 1 and 6 example prepns. of intermediates are included. In 1: Q is shown as II (dotted line may be a bond), R is C1-6alkoy, C1-6alkyl, C3-7cycloalkyl, Ph, and heteroaryl; T is piperarine-1,4-diyl; U is NR7, O, or S; V is C(H)kR1, O or N(R7)k; W

is

CR3 or NR10; X is CR4 or NR10; Y is CR5 or NR10; Z is CR6 or NR10; k is O or 1; m, n, and p are 0-2 provided that the sum of m, n, and p must equal 1 or 2; R8 and R8 are H, hydroxy, Cl-6alkyl, Cl-6alkoxy, cyano, and fluoro, or R8 and R8' taken together form :0, 15, 1NOR9, or 1NH; other variables and provisos are given in the claims.

474012-42-5P, 3-12-(4-Henroxylpiperazim-l-yl)-2-oxoethyl]-4-fluoro-1H-indole-7-carboxylic acid methylamide
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)

(drug candidate; preparation of indole, azaindole, and related heterocyclic

plperazinecarboxamides for treatment of AIDS)

ocyclic piperazinecarboxamides for treatment of AIDS) 474012-42-5 CAPLUS IH-Indole-7-carboxamide, 3-[2-(4-benzoyl-1-piperazinyl)-2-oxoethyl]-4-fluoro-N-methyl- (CA INDEX NAME)

L9 ANSWER 13 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2002:832569 CAPLUS

DOCUMENT NUMBER: 137:337880

137:337880
Preparation of indole, azaindole, and related heterocyclic piperazinecarboxamides for treatment of TITLE:

AIDS
Wang, Tao; Wallace, Owen B.; Meanwell, Nicholas A.;
Zhang, Zhongxing; Bender, John A.; Kadow, John F.;
Yeung, Kap-Sun
Bristol-Myers Squibb Company, USA
PCT Int. Appl., 111 pp.
CODEN: PIXXD2
Patent
English 1
1 INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM, COUNT: PATENT INFORMATION:

	TENT :																
WO	2002	0853	01		A2		2002	1031		WO 2	002-	US12	856			20020	423
WO	2002	0853	01		A3		2003	0227									
	W:															CH,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD	GE,	GH
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC	LK,	LR
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ	OM,	PH
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR	TT,	TZ
							ZA,										
	RW:	GH,	GM_{r}	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT	, BE,	CH
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	2445																
	2002																
	1381																
	R:												LU,	NL,	SE	MC,	PT
							RO,										
BR	2002	0091	53		A		2004	0720		BR 2	002-	9153				20020	423
CN	1520 1330	295			A		2004	0811		CN 2	002-	8126	29			20020	423
CN	1330	307			C		2007	0808									
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	2004																
	2003										003-					20031	
	2007				A1		2007	1220			007-					20071	
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										AU 2	002-	3075	05		АЗ	20020	423
										wn o	002-	rrs 1 2	856		w	20020	423

ANSWER 13 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

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ANSWER 14 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 2002:113840 CAPLUS
ACCESSION NUMBER:
                                              136:167283
Preparation of acetylpiperidinebutanediamines as calcium ion-permeable AMPA receptor antagonists Mimura, Tetsuya; Kawajiri, Shinichi Dalichi Selyaku Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 93 pp.
CODEN: JKXXAF
Patent
DOCUMENT NUMBER:
TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC, NUM. COUNT:
PATENT INFORMATION:
        PATENT NO.
                                              KIND DATE
                                                                                  APPLICATION NO.
                                                                                                                             DATE
JP 2002047272
PRIORITY APPLN. INFO.:
                                                           20020212
                                                                                  JP 2000-225300
JP 2000-225300
OTHER SOURCE(S):
                                              MARPAT 136:167283
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$$R^{1-}X-G^{N}$$
 $CH_{2}-A-Y-Q$

The compds. I (R1 = aryl, arylcarbonyl, aryloxy, cycloalkyl heterocyclyl, etc.; X = single bond, (un)substituted alkyl, alkenyl, cycloalkyl, monocyclic heterocyclyl; G = CO, SO2; n = 0-3; h = NR2, O, S, single bond;

monocyclic heterocyclyl G = O, SO2; h = D-3; A = NRZ, O, S, single bond;
R2 = H, alkyl, OH; Y = alkylene, alkynylene, alkenylene; Q = NR3R4, OR5, SR5; R3, R4 = H, alkyl, cycloalkyl, azalkyl, etc.; R5 = alkyl, cycloalkyl, azyl, heterocyclyl, etc.), their salts, and solvates are prepared The compds. are useful for cerebral infarction, senile dementia, Altheimer's, disease, Parkinson's disease, and Huntington's disease. Cycloalwal areacted with with oxalyl chloride in the presence of DMSO and Et3N in CH2Cl2 at -78° for 30 min and reacted with
4-[N-(4-aminobutyl)-N-(tert-butoxycarbonyl)aminomethyl]-1-(1-naphthylacetyl)piperidine for 1 h to give 828 N-(tert-butoxycarbonyl)-N-(1-(1-naphthylacetyl)piperidin-4-ylmethyl-1, 4-butanediamine, which was treated with HCl in EtcH at room temperature for 5 h to give N-cyclohexylmethyl-N'-[1-(1-naphthylacetyl)piperidin-4-ylmethyl-1, 4-butanediamine hydrochloride showing good AMPA receptor blocking activity in vitro

in vitro. 396071-91-3P 396071-92-4P SPOV.1-71-75 350V/1-72-48 RI. PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

L9 ANSWER 15 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2001:868447 CAPLUS
DOCUMENT NUMBER: 136:5917
TITLE: Preparation of (hetero)arylacyl-piperidinyl-benzylamines for use as tryptase inhibitors
INVENTOR(S): Astles, Peter C.; Eastwood, Paul R.; Houille, Ollvier:

Levell, Julian; Pauls, Heinz; Czekaj, Mark; Liang, Guyan; Gong, Yong; Pribish, James; Neuenschwander, Kent Aventis Pharmaceuticals Products Inc., USA PCT Int. Appl., 267 pp. CODEN: PIXXD2 Patent English 1 PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. WO 2001090101																
W :																
					21,	SL,	10,	1171,	IK,	, 11,	12,	UA,	06,	05,	02,	VIV,
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2003	0187	020	00,	A1	OP1	2003	1002	on,	IIS :	2001-	8431	26	10,	200	0010	426
6977	263			B2		2005	1220		-					_	0010	120
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1296	972			Al		2003	0402		EP	2001-	9309	25		2	0010	427
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20021	0056	01		A		2003	0106		NO :	2002-	5601			2	0021	121
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1057	899			A1		2006	0728		HK .	2004-	1007	65		2	0040	206
2005	0228	018		Al		2005	1013		US :	2005-	5780	9		. 2	0050	214
: APP	LN.	TNEO	. :						GB :	2000-	1236	2		A 2	UU 00	522
										2001-						
	RW: 2003 6977 2409 1296 R: 2001 2003 2004 1230 12002 2002 2002 2002 8586 605	CR, HUT, LUT, SD, FHC, CR, CR, CR, CR, CR, CR, CR, CR, CR, C	CR, CU, HU, ID, LU, LV, SP, SE, SP, SE, SP, SE, SE, SE, SE, SE, SE, SE, SE, SE, SE	CR, CU, CZ, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH	CR, CU, CZ, DE, HU, ID, IL, IM, LU, LV, MA, MD, SD, SE, SG, SI, YU, ZA, ZW RW: GH, GM, KE, LS, DE, DK, ES, FI, 20030187020 A1 1296972 A1 1296972 A1 1296972 A1 1296972 A1 1296972 A1 1296972 A1 1296971 A1 129030014891 A2 120030014891 A2 120030014891 A2 1200201501892 A2 1200201501892 A2 12002001484 A2 12002001484 A2 12002001484 A2 12002001484 A2 12002001484 A2 120020014899 A2 12002001484 A2 12002001484 A2 120020014889 A2 120020014889 A3 120030014889 A3 120030014881 A2 120020014881 A2 120020	CK, CU, CZ, DE, DK, H, H, L,	CR, CU, CZ, DE, DK, DM, MM, HU, ID, IL, IN, IS, JP, LU, LV, MA, MD, MG, MK, SD, SE, SG, SI, SK, SL, YU, ZA, ZW RN: GH, GM, KE, LS, MM, MZ, DE, DK, ES, FI, FR, GB, BJ, CF, CG, CI, CM, GR, 20030187020 Al 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20031129 CR AT, BE, CH, DE, DK, ES, FR, GB, GR, TT, LI, LV, NL, NL, FT, EJ, SI, LT, LV, FT, RO, MK, CY, AL, TR 200301285 A2 20031229 2004010897 T 20040408 2003002485 A3 20070928 2004510697 T 20040408 A 20030517 CN 2001-24098 A 2003017 A 2003017 A 2001-25017 CN 2001-27413 B2 20070118 A 20030521 A 20030521 A 20030523 MX 2002-11400 A 20030521 A 20030521 TAIGE9 A 20030521 TAIGE9 A 20030523 MX 2002-11400 A 20030521 TAIGE9 A 20030523 MX 2002-11400 A 20030521 TAIGE9 A 20030523 MX 2002-115683 B1 2008077 RX 2002-115683 B1 2008077 RX 2002-15683 B1 2008077 RX 2002-15683 BK 2002-115683 BK 2002-115683 BK 2002-15683	CR, CU, CZ, DE, DK, DM, DZ, DE, ES, FI, GB, GD, GE, GH, HU, DD, IL, IN, 1S, JP, KE, KG, KP, KP, KR, KZ, LC, LK, LK, LK, LU, LV, MA, MD, MG, MK, MN, MM, MX, MZ, ND, NZ, PL, PT, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, YU, ZA, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SD, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SD, 20030187020 Al 20030187020 Al 20031020 BZ 20051220 2409827 Al 20031020 BZ 20051220 2409827 Al 20031020 BZ 20051220 2409827 Al 200310402 BF 2001-843126 CA 2001-2409827 Al 200301402 BF 2001-843126 CA 2003-2409827 AL 200301402 BF 2001-2409827 AL 200301402 BF 2001-2409827 AL 200301405 AD 200302485 AD 200302485 AD 200302485 AD 200302485 AD 200302485 AD 2004028 AD 2001-2506014 AD 200300517 CM 2001-819152 AD 2001-2506014 AD 200300517 AD 2001-819152 AD 20000484 AD 20030231 AD 2001-81952 AD 2001-81958 AD 2001-81952 AD 2001-8	RM1 GH, GM, KE, LS, MW, MC, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, TE, TI, LU, MC, NI, FT, SE, CH, ROSON CH, SE, FIL, FR, GB, GR, TE, TI, LU, MC, NIL, FT, SE, CH, ROSON CH, SE, FIL, FR, GB, GR, TE, TI, LU, MC, NIL, FT, SE, CH, ROSON CH, SE, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH

CN 2001-811952

WO 2001-US13811

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L9 ANSWER 14 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) (prepn. of acetylpiperidinebutanediamines as calcium ion-permeable
      receptor antagonists)
396071-91-3 CAPLUS
Ethanone, 2-(5-fluoro-IH-indo1-3-y1)-1-[4-[[4-[(4-piperidinylnethyl)amino]butyl]amino]methyl]-1-piperidinyl]-,
rochloride
AMPA
hydr
                        (CA INDEX NAME)
                                                              СH2-NH- (СH2) 4-NH-СH2
          396071-92-4 CAPLUS Ethanone, 2-(5-fluoro-lH-indol-3-y1)-1-[4-[[4-[[(2S)-2-pyrsolidinylmethyl]amino]butyl]amino]methyl]-1-piperidinyl]-, hydrochloride (l:3) (CA INDEX NAME)
Absolute stereochemistry.
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●3 HCl

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ANSWER 15 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN R SOURCE(S): MARPAT 136:5917
                                                                                                                                                                                                                                                   (Continued)
    OTHER SOURCE (S):
    * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
   AB Title compds. I [Ar = (hetero)aryl, where the two groups on the Ar ring are \beta to each other; Rl-2 = H, alkyl; R3 = (un)substituted(hetero)aryl, arylalkenyl, cycloalkenyl, cycloalkyl, etc.; R4 = H, acyl, alkoxy, alkyloxycarbonyl, carboxy, CN, halo, etc.; n = 0 - 4] were prepared Over 300 synthetic examples were disclosed. For
R4 = H, acyl, alkoxy, alkyloxycarbonyl, carboxy, CN, halo, etc.; n = 0 - 4] were prepared Over 300 synthetic examples were disclosed. For instance,

3-bromobenzylbromide was converted in two steps to boronate II. II was coupled to the triflate ester derivative of the enol of 4-oxo-N-benzyloxycarbonylpiperidine (DMF, K2CO3, FdCl2(dppf)-CH2Cl2, 80°C, 18 h) to give the corresponding bicyclic intermediate. This intermediate was deprotected and reduced to the piperidine (EtOH, 10% Pd-C/H2, room temperature, 5 h) and coupled to

5-phenethylthiophene-2-carboxylic acid (DMF, HAPyU, iPr2NEt, room temperature, 18 h) to give III. III had Ki = 50

nM for tryptase. I are useful in the treatment of e.g., asthma and inflammatory diseases.

IT 375851-79-99

FL BSU (Biological study, unclassified); FAC (Pharmacological activity); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug; preparation of (hetero)arylacyl-piperidinyl-benzylamines for use as tryptase inhibitors)
                 tryptase inhibitors)
375851-79-9 CAPLUS
   NN 375017739 CAFLOS

(CN Ethanone, ethyl)phenyl]-1-piperidinyl]-2-(5-bromo-1H-indol-3-y1)-, 2,2,2-trifiluoroacetate (1:7) (CA INDEX NAME)
                       CM 1
                       CRN 375851-78-8
CMF C22 H24 Br N3 O
                       CM 2
```

A3 20010427

W 20010427

ANSWER 15 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L9 ANSWER 16 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2001:851126 CAPLUS DOCUMENT NUMBER: 135:371760
Preparation of pyrazolylpyrimidines and analogs as TNR-α signaling modulators
Sneddon, Scott F.; Kame, John L.; Hirth, Bradford H.;
Vinick, Fredy Qiao, Shuang; Nahill, Sharon R.
Genzyme Corporation, USA
PCT Int. Appl., 108 pp.
CODEN: PIXXD2 135:371760 TITLE: INVENTOR (S): PATENT ASSIGNEE(S): DOCUMENT TYPE: LANGHAGE .

English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 2001087849

W1 AB, AG, AL,

W1 CO, CR, CU,

GM, HR, HU,

LS, LT, LU,

RC, RU, SD,

UZ, VN, YU,

BF, DK, ES,

BJ, CF, CG,

CA 2408488

US 20020119988

US 6969728

EP 1294699

RI AT, BE, CH, A2 AM, CZ, ID, LV, SE, ZA, LS, FI, CI, A1 A1 B2 A2 BR 2001011158 BR 2001011158 AU 2001259691 MX 2002010993 NO 2002005405 NO 324693 KR 840816 US 20040171617 US 7034031 20071203 KR 2002-715152 US 2004-797244 20021112 20080623 A1 20040902 20040310 20060425 US 20060173010 US 2005-292325 20051201 20060803 PRIORITY APPLN. INFO.: US 2000-203784P P 20000512 US 2000-205213P P 20000518 IIS 2001-852965 A3 20010510 WO 2001-US15027 W 20010510

OTHER SOURCE(S): MARPAT 135:371760

(Continued) ANSWER 16 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN

Title compds. [I; R1 = H or NH2; R2 = ZZ3(CH2)nR; R = (un)substituted Ph or -heterocyclyl; R4 = (alky1-substituted) 2-pyridinyl or -pyrazinyl; Z = (un)substituted pyrazole-1,4-diy1; Z1,Z2 = N or CH; Z3 = O, CH2, S, SO2;

= 0-2] were prepared Thus, 4-(Me2HC)C6H4OH was condensed with (MeCO) 2CHN2

(MecO) ZCHN2
and the product cyclocondensed with
4-(2-pyridinyl)-2-pyrimidinylhydrazine
to give title compound II. Data for biol. activity of I were given.
IT 374080-55-4P 374080-62-3P

RL: BAC (Biological activity or effector, except adverse); BSU

(Biological

modulators)
374080-55-4 CAPLUS
1H-Indole-3-acetamide, 5-bromo-N-[1-(3-cyanopheny1)-2-[[2-(4-methoxypheny1)ethy1]amino]-2-oxoethy1]-N-[2-(1H-imidazo1-5-y1)ethy1]-

INDEX NAME)

374080-62-3 CAPLUS
1H-Indole-3-acetamide, 5-bromo-N-[1-(3-cyanophenyl)-2-[(2,2-diphenylethyl)amino]-2-oxoethyl]-N-[2-(1H-imidazol-5-yl)ethyl]- (CA

ANSWER 16 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

US 2004-797244

A1 20040310

02/02/2009 10/539,151

L9 ANSWER 17 OF 44 C	APLUS (COPYRIGHT 20	09 ACS on STN	
ACCESSION NUMBER:	2001:	762989 CAPL	US	
DOCUMENT NUMBER:	135:3	18419		
TITLE:	Synthe	esis of subs	tituted bipiperidine:	s and their use
		antagonists		
INVENTOR(S):			Rigby, Aaron; Sangan	nee, Hitesh;
		gthorpe, Bri		
PATENT ASSIGNEE(S):		zeneca AB, S		
SOURCE:		nt. Appl., 1	60 pp.	
		: PIXXD2		
DOCUMENT TYPE:	Patent			
LANGUAGE:	Englia	n.		
FAMILY ACC. NUM. COUNT:	7			
PATENT INFORMATION:				
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FAILNI NO.			AFFEIGRIION NO.	
WO 2001077101	A1	20011018	WO 2001-SE751	20010405

	TENT NO.																
											2001-						
	₩:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE	, ES,	FI,	GB,	GD,	GE,	GH,	GM,
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		LT,	LU,	LV,	MA.	MD,	MG,	MK,	MN,	MW	, MX,	MZ,	NO,	NZ,	PL,	PT,	RO,
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		BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML	, MR,	NE,	SN,	TD,	TG		
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EΡ	1274	701			B1		2005	0629									
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CN	1244	576			C		2006	0308									
JP	2003	5303	93		T		2003	1014		JP	2001-	5755	74		2	0010	405
NZ	5215	43			A		2004	1029		NZ	2001-	5215	43		2	0010	405
ΕP	1493	743			A1		2005	0105		EP	2004-	2059	9		2	0010	405
ΕP	1493	743			B1		2008	0903									
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CN	1660	839			A		2005	0831		CN	2004-	1010	2245		2	0010	405
ΑU	2001	2469	97		B2		2007	0329		ΑU	2001-	2469	97		2	0010	405
ΑT	4071	31			T		2008	0915		AT	2004-	2059	9		2	0010	405
US	2002	0077	337		A1		2002	0620		US	2001-	8274	88		2	0010	406
US	6525	070			B2		2003	0225			2001- 2004- 2001- 2001-						
ZA	2002	0077	00		A		2004	0102		ZA	2002-	7700			2	0020	925
NO	2002	0047	74		A		2002	1129		NO	2002-	4774			2	0021	003
MΧ	2002	0098	85		A		2003	0327		MX	2002-	9885			2	0021	007
US	2004	0006	080		A1		2004	0108		US	2003-	3410	27		2	0030	113
US	6903	115			B2		2005	0607									
US	2004	0014	783		A1		2004	0122		US	2003-	4365	82		2	0030	513
US	7238	811			B2		2007	0703									
ΗK	1051	193			A1		2005	1028		HK	2002- 2002- 2002- 2003- 2003- 2003-	1034	24		2	0030	514

L9 ANSWER 17 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) thioheterocycly1) were prepd. Examples include: data for over 600 compds., 4 solid oral dosage and 1 parenteral (general) formulations, a bloassay for Ca2+ flux, human eosinophil chemotaxis and H1 antagonism. E.g., 4 (3, 4-dichlorophenoxy)piperidine was alkylated with 1-(tert-butoxycarbonyl)-4-piperidone (1,2-dichloropethane, NaBH(OAc), HOAC, 18 h, room temp.) to give an intermediate [1,4']bipiperidine. This intermediate was deprotected (DCM, TFA, 4 h, room temp.) and the resulting bipiperidine condensed with 3-methanesulfonylbenzoic acid (THF, PYBROP, (i-PP)ZNET, 18 h, room temp.) to give example compd. II isolated as the acetate salt. I are used in the treatment of a chemokine (such as CCR3) or H1 mediated disease state.

IT 367497-01-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study) PREP (Preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (drug; synthesis of substituted bipiperidines and use as H1 antagoniats)

RN 367497-01-6 CAPLUS

Ethanone,
1-[4-(3,4-dichlorophenoxy) | 1,4'-bipiperidin]-1'-y1]-2-(5-hydroxy-1H-indol-3-y1)- (CA INDEX NAME)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9	ANSWER 17 OF US 2005017109 US 7179922		COPYRIGHT 2009 20050804 U 20070220	ACS on STN S 2005-76773	(Conti	nued) 20050310
PRIO	US 2007017929 RITY APPLN. IN	7 A1	20070802 U	S 2007-732411 B 2000-8626	A	20070403
			G	в 2000-19111	A	20000803
			S	E 2000-3664	Α	20001011
			C	N 2001-810683	EA.	20010405
			E	P 2001-920053	A3	20010405
			W	O 2001-SE751	W	20010405
			σ	S 2001-827488	EΑ	20010406
			ū	S 2003-341027	A1	20030113
			U	S 2003-436582	A3	20030513

MARPAT 135:318419

(CH₂)_n— (CH_Y)_q— (CH₂)_r—R³

OTHER SOURCE(S):

Title compds. I |q, s, t=0-1; n, r=0-5; m, p=0-2; X=CH, C(O), O, S, S(O), S(O), N-; provided that when m and p are both 1 then X is not CH; Y=NHR2, OH; T=C(O), C(S), S(O), CH2; R1=H, alkyl, aryl, heterocyclyl; R2, R47=H, alkyl, aryl-alkyl, CO-alkyl; R3=alkyl, alkenyl, cycloalkyl, cycloalkenyl, aryl, heterocyclyl, thioaryl, AB

L9 ANSWER 18 OF 44
ACCESSION NOMBER:
DOCUMENT NUMBER:
135:303899

Synthesis of heterocycloalkylbenzocyclobutanes and heteroarylbenzocyclobutanes and their use as inhibitors of serotonin and noradranilm reuptake
PATENT ASSIGNEE(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:
FATENT INFORMATION:
1

2001:760046 CAFUUS
Coptuments of heterocycloalkylbenzocyclobutanes and their use as inhibitors of serotonin and noradranilm reuptake
Bettrand, Millan, Mark; Lejeune, Francoise; Brocco,
Mauricette
Addr Et Compagnie, Fr.; Servier Lab
EU: Pat. Appl., 47 pp.
COODN: EPXXDW
French
FAMILY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:
FATENT INFORMATION:
1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

		NO.				DATE									DATE	
						2001									20010	412
EP	1146	041			B1	2003	1112		200		.005					
						ES,		GI	R. I	IT.	LI.	LU.	NL,	SE.	MC.	PT.
						RO										
FR	2807	753			A1	2001	1019	FR	200	00-4	1742			- 2	20000	413
FR	2807	753			В1	2002	0607									
MX	2001	0035	53		A	2002	0604	MΧ	200	01-3	3553			- 2	20010	406
JP	2001	3025	99		A	2001	1031	JP	200	1-1	111	69		- 2	20010	410
JP	3761	796			B2	2006	0329									
NO	2001	0018	62		A	2001	1015	NO	200	11-1	862			2	20010	411
						2005										
		0014				2001									20010	
		0030				2001									20010	
		0019				2002		US	200	01-8	338	27		- 2	20010	412
		413				2002										
		0015				2002		HU	200	01-1	.503			- 2	20010	412
		0015				2003										
		92				2002									20010	
		02				2003									20010	
		041			T	2004									20010	
		104				2004										
		25													20010	
		794			A	2001		CM	200	01-1	.163	86		- 2	20010	413
		659				2004										
		255				2001		CA	200	01-2	2344	255		- 2	20010	417
		255				2006	0711									
		477			A1	2005	0506								20020	
CORIT:	' APF	LN.	INFO	. :				FR	200	00-4	1742		-	A 2	20000	413

OTHER SOURCE(S): MARPAT 135:303899 L9 ANSWER 18 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

-(CH₂)_n-N-R1

AB Title compds. I [n = 1 - 6; Rl-2 = H, alkyl, aryl, arylalkyl, cycloalkyl(alkyl), alkenyl, alkynyl, heterocyclyl, etc.; X = CH:CH, O, SOO-2, NR3; Y = CH/CH2; T = cycloalkyl (mono or polycyclic), heterocyclyl)

were prepared Forty example compds. were disclosed. E.g., 6-cyano-1-methylsulfonyl-5,6-dihydrocyclobuta[f]indole (preparation given) was desulfonylated (N Noov - CO)

desulfonylated (K, MeOH, reflux, 12 h) and converted to tetrahydro

derivative
II (HOAc, NaCNBH3, room temperature, 2 h). II was alkylated with

hexanone
(THF, n-BuLi, -80°C) and the resulting nitrile reduced to
aninomethyl derivative III (MeOB, B2-Ra/Ni, 30 bar, 60°C, 24 h). In
competitive binding assays, compds. of the invention showed affinity for
serotonin reuptake binding sites, pXi > 7 and noradenaline reuptake
binding sites, pXi ≥ 6. I are used to treat depression, panic
attacks, anxiety, obesity, etc.
367263-60-3P

367263-60-3P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate; synthesis of heterocycloalkylbenzocyclobutanes and heteroarylbenzocyclobutanes and their use as inhibitors of serotonin and noradrenaline reuptake) 367263-60-3 CAPLUS 1H-Indole-3-acetamide, N-[[1,2-dihydro-1-(1-

hydroxycyclopenty1)cyclobuta[b]naphthalen-1-y1]methy1]-5-fluoro-N-methy1-(CA_TNDEX_NAME)

ANSWER 18 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

-OH

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L9 ANSWER 19 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
136:179
From Hit to Lead. Combining Two Complementary Methods for Focused Library Design. Application to μ Opiate Ligands
AUTHOR(S):

CORPORATE SOURCE:
SOURCE:
Department of Chemistry, CEREP, Lille, F-59000, Fr.
Journal of Medicinal Chemistry (2001), 44(21), 3378-3390
CODEN: JOURNARY, ISSN: 0022-2623
American Chemical Society
Journal
LANGUAGE:
CHER SOURCE(S):
CASREACT 136:179

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

Compound I obtained by random screening and displaying a micromolar

AB Compound I obtained by random screening and displaying a micromolar activity

on the μ opiate receptor was chosen as a starting point for optimization. Two complementary concepts of similarity were used for the design of analogs and compared. These are based, resp., on a computer-aided comparison of pharmacophoric patterns and on topol. similarity. The structure-activity relationships are discussed in light of both similarity concepts. An N-methyl-3-(4-oxo-1-phenyl-1,3,8-triaraspiro(4.5)decyl)acetamide derivative, designed by combining the structure-activity relationships enlightened by each method, has a subnanomolar affinity for μ (h) receptor (ICSO = 0.9 mM). It is a promising lead, allowing the design of a new series of analogs substituted at the N-3 of the spirocycle moiety.

IT 372956-13-3P RL: RCT (Reactant); SFN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (combining two complementary methods for focused library design and application to μ opiate ligands)

RN 372956-13-3 CAPLUS

Ethanone, 2-(5-bromo-1H-indol-3-yl)-1-[4-(7-nitro-2,1,3-benzoxadiazol-4-yl)-1-piperazinyl]- (CA INDEX NAME)

ANSWER 19 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

REFERENCE COUNT: THERE ARE 26 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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ANSWER 20 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
 ACCESSION NUMBER:
                                                                                         2001:662562 CAPLUS
  DOCUMENT NUMBER:
                                                                                         135:352346
 TITLE:

From Hit to Lead. Analyzing Structure-Profile
Relationships
AUTHOR(S):

Poulain, Rebecca; Horvath, Dragos; Bonnet, Beatrice;
Eckhoff, Christian; Chapelain, Beatrice; Bodinier,
Marie-Christine; Peprez, Benoit
CORPORATE SOURCE:

Department of Chemistry, CRREP, Lille, F-59000, Fr.
JOURNAL of Medicinal Chemistry (2001), 44(21),
3391-3401

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER:
DOCUMENT TYPE:
Journal
LANGUAGE:

American Chemical Society
Journal
LANGUAGE:
English
AB Two compds., (piperidine and piperazine carboxylic acid derivs.) obtained
by random screening, and displaying micromolar activities on the µ
opiate receptor were used as starting points for optimization. In that
work, the traditional concept of the activity of a compound (related to one
                                                                                          From Hit to Lead. Analyzing Structure-Profile
  TITLE:
PUBLISHER:
                   or a few targets) was extended to the comprehensive pharmacol. profile of that compound on more than 70 receptors, transporters, and channels
that compound on more than 10 receptors, compositions to a CNS-oriented project. Using the two complementary design strategies based on two similarity concepts described in the previous paper, we have obtained analogs with IC50 values ranging between 0.9 mW and a few micromolar on the µ receptor and displaying qual. different profiles. We discuss here, both on a case-by-case basis and from a statistical standpoint, the pharmacol. profiles in light of the two similarity concepts.
                    concepts.
372956-13-3
RL: BAC (Biological activity or effector, except adverse); BSU
  (Biological
                  logical study, unclassified); BIOL (Biological study) (piperidine- and piperazine carboxylic acid derivative opioid receptor structure-activity relationship, and compound preparation) 372956-13-3 CAPLUS Ethanone, 2-(5-bromo-1H-indol-3-yl)-1-[4-(7-nitro-2,1,3-benzoxadiazol-4-yl)-1-piperazinyl] (CA INDEX NAME)
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REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR

L9 ANSWER 21 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2001:565002 CAPLUS
DOCUMENT NUMBER: 135:152713
Aromatic amides as novel melanocortin receptor agonists and antagonists
INVENTOR(S): Lundstedt, Tozbjezn; Skottner, Anna; Seifert, Elisabeth; Starchenkov, Igor; Trapencleris, Peteris; Kauss, Valerjans, Kalvins, Ivars; Boman, Arne Melacure Therapeutics AB, Swed.
PCT Int. Appl., 52 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: PRINTACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

														NO.				
	WO		0551	06		A2		2001	0802					46				
		W:	AE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BE	3, BG	, BR	, BY,	BZ,	CA,	CH,	CN,
			CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES	, FI	, GB	, GD,	GE,	GH,	GM,	HR,
			HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KF	, KF	, KZ	, LC,	LK,	LR,	LS,	LT,
			LU,	LV,	MA,	MD,	MG,	MK,	MN,	ΜW,	MX	, M2	, NO	, NZ,	PL,	PT,	RO,	RU,
			SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TF	, TI	, TZ	, UA,	UG,	US,	UZ,	VN,
			YU,	ZA,	ZW													
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, T2	, UG	, ZW,	AT,	BE,	CH,	CY,
														, NL,				BF,
														, SN,				
	CA	2398	728			A1		2001	0802		CA	2001	-239	8728		2	0010	129
														3				
	EP													850				
		R:												, LU,	NL,	SE,	MC,	PT,
								RO,										
														048				
	ZA	2002	0058	86		A		2004	0621		ZA	2002	-588	6 9		2		
																	0020	
DD 701								2003	1016					192 8				
PRIO	KT.T.7	APP	LN.	TNFO	. :						GB	2000	-194	8		A 2	0000	178
											GB	2000	-206	0		A 2	0000	128
											WO	2001	-GB3	46		w 2	0010	129
OTHE	3 80	DURCE	(S):			MAR:	PAT	135:	1527.	13								

The present invention relates to novel aromatic amides (1; B-E-X-N(R8)-C(O)-Y-F-A and pharmacol. active salts thereof) and to the

of these amides for the treatment of obesity, anorexia, inflammation, mental disorders and other diseases associated with the melanocortin receptors or related systems, e.g. the melanocyte stimulating hormones. In I. E and F are independently a saturated or unsatd., acyclic hydrocarbon

group having 1-5 C atoms. X and Y are independently methylene; one of X and Y are absent (i.e. a single bond); or X can be -CH(QR10)- and/or Y

be -CH(MR9)- (M and Q are independently a saturated or unsatd., straight branched chain acyclic hydrocarbon group with 1-6 C atoms; or M and/or Q L9 ANSWER 20 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

10/539,151

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L9 ANSWER 21 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) are absent (i.e. M and/or Q are single bonds)). R8, R9 and R10 are H, -PR4, -C(0)DR4 (P and D are independently a satd. or unsatd, straight or branched chain acyclic hydrocarbon group having 1-6 C atoms; or D is a beent (i.e. D is a single bond)). R4 is hydroxy, Me, cyclohexyl, cyclopentyl, aminoquanidine, guarbodine, carboxy, or (possibly substituted) amino, carbamoyl, alkoxy, alkoxycarbonyl, acyl, morpholinyl, pyrrolidinyl, piperidinyl, pherazinyl, Ph, isoindolyl, indenyl, pyridinyl, indolyl, pyrrolyl, cyclopentadienyl wherein R4 in R8, R9 and R10 may be the same or
```

pyrroly1, cyclopentadienyl wherein R4 in R8, R9 and R10 may be the same different. A and B are the same or different and are (possibly substituted) quinolinyl, isoquinolinyl, isoquinolyl, ngathyl, pyridinyl, indolyl, pyrazinyl, cyclopentadienyl, pyrimidinyl, Fh, indenyl. Several claimed compds. (N-(3-aminopropyl)-3-(1H-indol-3-yl)-2-(2-naphthalen-1-ylacetylamino)propionamide hydrochloride (1:1.2), N-(1-[benzyl(4-quanidinobutyl)carbamoyl]-2-(1H-indol-3-yl)-2+(2-naphthalen-1-yhenylbutyramide monohydrochloride,
N-benzyl-N-(4-quanidinobutyl)-3-(1H-indol-3-yl)-2-(2-naphthalen-2-ylacetylamino)propionamide monohydrochloride,
N-[1-(9-ethyl-9H-carbazol-3-ylcarbamoyl)-2-(1H-indol-3-yl)-thyl]-4-quanidinobutyramide monohydrochloride,
4-amino-N-[1-(9-ethyl-9H-carbazol-3-ylcarbamoyl)-2-(1H-indol-3-yl)-thyl)-pylbutyramide monohydrochloride,
2-(3-aminopropionylamino)-N-(9-ethyl-9H-carbazol-3-yl)-3-(1H-indol-3-yl)-propionanide monohydrochloride) were tested (zesults given) for affinity for melanocortin receptors (MCL, MC3, MC4, MC5) and/or influence on cAMP. In vivo effects on food intake and anti-inflammatory effects were also detd. on selected compds. Two example prepns. are given.

RLI BAC (Biological activity or effector, except adverse); BSU

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

(aromatic animes __
antagonists
and their preparation)
RN 352277-28-2 CAPLUS
CN 1H-1ndole-3-acetamide,
5-bromo-N-[1-(2-bromopheny1)-2-(cyclohexylamino)-2oxoethyl]-N-[2-(dimethylamino)ethyl]-, hydrochloride (1:1) (CA INDEX NAME)

L9 ANSWER 21 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L9 ANSWER 22 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 2001:83714 CAPLUS DOCUMENT NUMBER: 134:311061

Synthesis of 5-(sulfamovlmethyl)indoles TITLE: AUTHOR(S):

Bosch, J.; Roca, T.; Armengol, M.; Fernandez-Forner,

AUTHOR(S):

Bosch, J.; Roca, T.; Armengol, M.; Fernandez-Forner, D.

CORPORATE SOURCE:

Laboratory of Organic Chemistry, Faculty of Pharmacy, University of Barcelona, Barcelona, 08028, Spain Tetrahedron (2001), 57(6), 1041-1048

CODEN: TETRAB; ISSN: 0040-4020

Elsevier Science Ltd.

DOUMENT TTPE:

DOUMENT TTPE:

Journal

AB The synthesis of 5-(sulfamoylmethyl)indoles bearing a two-carbon chain at C-3 (aminoethyl, acetate, hydroxyethyl, ethyl) either by the Grandberg modification of the Fischer indolization or by intramol. Heck reaction of suitable o-halotrifluoroacetanilides is reported.

IT 334961-21-4P

RL: RCT (Reactant); SFN (Synthetic preparation); FREF (Preparation); RACT (Reactant on Teagent)

(preparation of 5-(sulfamoylmethyl)indoles)

RN 334981-21-4 CAPLUS

NN 18-Indole-3-acetamide,
5-[[[(1,1-dimethylethyl)amino]sulfonyl]methyl]-N,N-dimethyl- (CA INDEX NAME)

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 23 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2000:762637 CAPLUS
DOCUMENT NUMBER: 134:86:116
TITLE: Design, Synthesis, and Biological Evaluation of

and Selective Amidino Bicyclic Factor Xa Inhibitors
Han, Qi, Dominguez, Celia; Stouten, Pieter F. W.;
Park, Jeongsook M.; Duffy, Daniel E.; Galemmo, Robert
A., Jr.; Rossi, Karen A.; Alexander, Richard S.;
Smallwood, Angela M.; Wong, Pancras C.; Wright,
Matthew M.; Leuttegn, Joseph M.; Knabb, Robert M.;
Wexler, Ruth R.
DuPont Pharmaceuticals Company, Wilmington, DE,
19880-0500, USA
Journal of Medicinal Chemistry (2000), 43(23),
4388-4415
CODEN: JMCMAR; ISSN: 0022-2623
American Chemical Society
Journal AUTHOR(S):

CORPORATE SOURCE:

Journal English CASREACT 134:86116

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

A novel series of factor Xa (fXa) inhibitors incorporating an amidino 6,5-fused bicyclic moiety, e.g. I (R=Me, F, Cl, Br, etc.), has been designed and synthesized based on mol modeling studies. Structure-activity relationship (SAR) studies have led to selective subnanomolar fXa inhibitors. The most potent fXa inhibitor in this

series

I (R = Br) has a potent inhibition constant (Ki = 0.3 nM), is 350-fold selective for fXa over trypsin, and also shows good in vivo efficacy in rabbit atterio-venous thrombosis model (1D50 = 0.14 pmol/kg/h). An X-ray crystal structure of I (R = Br) complexed to bovine trypsin was completed, and its binding mode with fXa has been proposed based on modeling with human des-Gla-fXa.

IT 202124-24-IP RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic present)

inhibitors) 202124-24-1 CAPLUS

ANSWER 23 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) 1H-Indole-3-acetamide, 5-(aminoiminomethyl)-N-[2'-(aminosulfonyl)[1,1'-biphenyl]-4-yl]-N-methyl- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

316364-41-7P RL: RCT (Reactant); SFN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and antithrombotic activities of amidino bicyclic factor

Хa

Xa inhibitors)
RN 316364-41-7 CAPLUS
CN 1H-Indole-3-acetamide,
N-[2'-(aminosulfonyl) [1,1'-biphenyl]-4-yl]-5-cyanoN-methyl- (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 43 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 24 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2000:31350 CAPLUS 132:78470 DOCUMENT NUMBER:

Preparation of spiro-substituted azacycles as TITLE:

Preparation of approvements of the neurokinin antagonists
Maccoss, Malcolm; Mills, Sander G.; Shah, Shrenik K.;
Chiang, Yuan-ching P.; Dunn, Patrick T.; Koyama, INVENTOR(S):

Merck and Co., Inc., USA U.S., 49 pp. CODEN: USXXAM Patent PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION: English

PATENT NO. KIND DATE APPLICATION NO. DATE US 6013652 PRIORITY APPLN. INFO.; 19971204 19971204 20000111

OTHER SOURCE(S): MARPAT 132:78470

The title compds. [I, 1, m = 0-5 (with the proviso that l + m = 1-5); R1

H, alkyl, alkenyl, etc.; W = a bond, (un)substituted alkyl, Q = 0, S, S0, S02, NR2 (with the proviso that when W = a bond and X = alkyl, then Q

be NR2; R2 = H, alkyl, etc.); X = a bond, (un)substituted alkyl, NRCO, etc.; YZ considered together are 2 adjoining atoms of Ph, naphthyl, heteroaryl; the nitrogen in one of the rings is optionally quaternized with alkyl or phenylalkyl or is optionally present as an N-oxidel, tachykinin receptor antagonists useful in the treatment of inflammatory diseases, pain or migraine, and asthma, were prepared E.g., a 2-step synthesis of 3-(S)-II was given. In particular compds. I are shown to be neurokinin antagonists, and, e.g., they have been found to displace radioactive ligand for the NK-1 receptor at 0.01 nM to 1.0 μM , for the NK-2 receptor , 0.01 nM to 5 μM , and for the NK-3 receptor, 1.0 nM to 10 μM .

L9 ANSWER 25 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1999:635463 CAPLUS
DOCUMENT NUMBER: 131:243191
TITLE: Spiro-substituted azacycles as modulators of

TITLE: chemokine

receptor activity
Mills, Sander G.; MacCoss, Malcolm; Springer, Martin INVENTOR(S):

S.
Merck and Co., Inc., USA
U.S., 97 pp.
CODEN: USXXAM
Patent
English
2 PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO US 5962462 PRIORITY APPLN. INFO.: US 1997-989947 US 1996-32735P 19971212 19961213 19991005 US 1996-33558P P 19961220

OTHER SOURCE(S): MARPAT 131:243191

AB The invention is directed to spiro-substituted azacycles which are useful as modulators of chemokine receptor activity. Specifically, I [R1 = H, (un)substituted alk(en/yn)yl; W = bond, (un)substituted alkylene; O = (un)substituted alkylene, S, S(O), S(O), S(O), S(O), S(O), S(O), C(O), Etc.; YZ = fused aryl or heteroaryl nucleus; m, n = 0 to 5; (m+n) = 1 to 5] were prepared The compds. are useful as modulators of the chemokine receptors CCR-1, CCR-2, CCR-2A, CCR-2B, CCR-3, CCR-4, CCR-5, CXCR-3, and/or CXCR-4 (no data), and are thereby useful as antiinflammatory and immunomodulating agents. Use for the treatment of HIV infection and/or AIDS is claimed specifically. For instance, 1'-methylspiro[indoline-3,4'-piperidine] underwent a sequence of

ANSWER 24 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN 167485-09-8P(Continued) RL: BAC (Biological activity or effector, except adverse); BSU (Biological logical study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation of spiro-substituted azacycles as neurokinin antagonists) 167485-09-8 CAPLUS

16/485-U9-8 CAPLUS Ethanone, 2-dihydro-1-(methylsulfonyl)spiro[3H-indole-3,4'-piperidin]-1'-yl]-2-(5-fluoro-1H-indol-3-yl)- (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 26 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 25 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) N-benzoyloxycarbonylation (71%), N'-demethylation (73%), reductive N'-alkylation with a corresponding polyfunctional aldehyde, and removal

of
the benzoyloxycarbonyl protecting group, to give title compd. II.
IT 167485-09-8P
RI: BAC (Biological activity or effector, except adverse); BSU
(Biological
study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
(Reactant or reagent); USES (Uses)
(target compound; preparation of spiro-substituted azacycles as
modulators of

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

ANSWER 26 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN .SSION NUMBER: 1999:205361 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 130:252241

TITLE: Preparation of amidinoindoles and analogs as factor

inhibitors INVENTOR(S): Dominguez, Celia; Han, Qi; Duffy, Daniel Emmett;

Jeongsook Maria; Quan, Mimi Lifen; Rossi, Karen

Anita:

PATENT ASSIGNEE(S):

Wexler, Ruth Richmond Dupont Pharmaceuticals Company, USA U.S., 46 pp. CODEN: USXXAM Patent English 1 DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5886191	A	19990323	US 1997-916736	19970818
US 6043257	A	20000328	US 1998-176037	19981021
PRIORITY APPLN. INFO.:			US 1997-916736 A	3 19970818

MARPAT 130:252241 OTHER SOURCE(S):

AB

CO, CONH, etc.; Z1 = C6H4, CH2C6H4, pyridine-2,4-diyl, etc.; n = 0 or 1; dashed line = optional addnl. bond] were prepared as factor Xa inhibitors (no data). Thus, 5-cyanoindole was acylated by (COCl)2 and the product converted in 3 steps to 5-cyanoindole-3-acetic acid which was amidated by 4-(2-aminosulfonylphenyl)-2-pyridinamine to give, in 2 addnl. steps, I

ANSWER 26 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) (aminoiminomethyl)phenyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

202123-98-6 CAPLUS
Glydine, N-[[3-[aminoiminomethyl]]-1-methyl-1H-indol-3-yl]acetyl]-N-[[4[aminoiminomethyl]phenyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} Me & NH \\ N & CH_2-C-OMe \\ \end{array}$$

202124-01-4 CAPLUS

1H-Indole-5-carboximidamide, 1-methyl-3-[2-oxo-2-[4-(phenylmethyl)-1-piperazinyl]ethyl]- (CA INDEX NAME)

202124-04-7 CAPLUS 1H-Indole-5-carboximidamide, 3-[2-oxo-2-(4-phenyl-1-piperazinyl)ethyl]-(CA INDEX NAME)

202124-24-1 CAPLUS

ANSWER 26 OF 44 CAPLUS COFYRIGHT 2009 ACS on STN (Continu 202124-04-7P 202124-24-1P 202124-28-5P 202126-86-1P RL: BAC (Biological activity or effector, except adverse); ESU (Continued)

RI: BAC (Biological activity or elector, color----(Biological)
study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
(prepn. of amidinoindoles and analogs as factor Xa inhibitors)
RN 202123-90-8 CAPLUS
CN 1H-Indole-5-carboximidamide, 3-[2-[4-[(4-methylphenyl)sulfonyl]-1-piperazinyl]-2-oxoethyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

202123-94-2 CAPLUS
1H-Indole-5-carboximidamide, 3-[2-oxo-2-[4-[(phenylmethyl)sulfonyl]-1-plperazinyl]ethyl]- (CA INDEX NAME)

$$\underset{\mathbb{N}H}{\overset{H}{\longrightarrow}} \underset{\mathbb{N}H}{\overset{G}{\longrightarrow}} \underset{\mathbb{N}H}{\overset{\mathbb{N}H}{\overset{G}{\longrightarrow}} \underset{\mathbb{N}H}{\overset{G}{\longrightarrow}} \underset$$

202123-96-4 CAPLUS
1H-Indole-5-carboximidamide, 3-[2-oxo-2-[4-(phenylmethyl)-1-piperazinyl]ethyl]- (CA INDEX NAME)

$$\underset{\mathbb{N}H}{\text{H2N-CH2-Ph}}$$

ANSWER 26 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) lH-Indole-3-acetamide, 5-(aminoiminomethyl)-N-[2'-(aminosulfonyl)[1,1'-biphenyl]-4-yl]-N-methyl- (CA INDEX NAME)

202124-28-5 CAPLUS
1H-Indole-3-acetamide, 5-(aminoiminomethyl)-N-methyl-N-{2'[(methylamino)sulfonyl][1,1'-biphenyl]-4-yl]- (CA INDEX NAME)

202126-86-1 CAPLUS 1H-Indole-5-carboximidamide, 3-(2-oxo-2-[4-(phenylmethyl)-1-plperidinyl]ethyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

Searched by Jason M. Nolan, Ph.D.

ANSWER 26 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

● HCl

REFERENCE COUNT:

23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L9 ANSWER 27 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1999:96240 CAPLUS 130:153571
Preparation of indole and 2,3-dihydroindole derivatives as potent serotonin reuptake inhibitors and 5-HTIA receptor antagonists Moltzen, Ejner Knud; Perregaard, Jens Kristian; Mikkelsen, Ivan; Smith, Garrick Paul H. Lundbeck A/S, Den.
PCT Int. Appl., 47 pp.
CCOEN: PIXXD2
Patent
Forligh DOCUMENT NUMBER: TITLE: INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE. English DOCUMENT TIPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:

											LICAT						
											1998-						
											, BY,						
											, HU,						
		KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU	, LV,	MD,	MG,	MK,	MN,	MW,	MX
		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG	, SI,	SK,	SL,	TJ,	TM,	TR,	TI
		UA,	UG,	US,	UZ,	VN,	YU,	ZW									
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW	, AT,	BE,	CH,	CY,	DE,	DK,	ES
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL	, PT,	SE,	BF.	BJ,	CF.	CG,	CI
		CM	CA	CTAT	CTAT	BAT	BATO	BIR	CTAT	TT	TC						
ZA	9806	237			A		1999	0331		ZA	1998- 1998-	6237			1	9980	714
CA	2297	825			A1		1999	0204		CA	1998-	2297	825		1	9980	720
CA	2297	825			C		2006	0314									
AU	9885	340			A		1999	0216		TIA	1998-	8534	Ω		1	9980	720
ΑU	7365	96			B2		2001	0802									
ΕP	1007	523			A1		2000	0614		EP	1998-	9362	70		1	9980	720
ΕP	1007	523			B1		2003	1022			1998-						
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	P:
TR	2000	0023	1		T2		2000	0721		TR	2000-	231			1	9980	721
BR	9810	790			A		2000	0725		BR	1998-	1079	0		1	9980	721
ΗU	2000	0028	30		A2		2001	0928		ΗU	2000-	2830			1	9980	721
ΗU	2000	0028	30		A3		2001	1029									
HU	2251	01			В1		2006	0628									
NZ	5022	52			A		2001	0928		NZ	1998- 2000- 1998- 2000-	5 0 22	52		1	9980	721
JP	2003	5245	71		T		2003	0819		JΡ	2000-	5041	36		1	9980	720
CN	1127	501			C		2003	1112		CN	1998- 1998- 1998-	8075	54		1	9980	720
ΑT	2525	75			T		2003	1115		AT	1998-	9362	70		1	9980	720
PT	1007	523			T		2004	0227		PT	1998-	9362	70		1	9980	720
ES	2206	963			Т3		2004	0516		ES	1998-	9362	70		1	9980	720
					A		2004	0728		CN	2003-	2003	1060	02	1	9980	720
	1286				C		2006	1129									
	1515				A		2004	0728		CN	2003-	2003	1060	03	1	9980	720
	1293				C		2007	0103									
CZ	2959	37			В6		2005	1214		CZ	2000-	285			1	9980	72
					В6		2006	0105		SK	2000-	95			1	9980	720
	1909	24			D 1		2006	0000									

L9	ANSWER 27 OF 44	CAPLUS	COPYRIGHT 200			(Contin	
	IN 1998MA01631	A	20050304	IN	1998-MA1631		19980722
	MX 2000000700	A	20010131	MX	2000-700		20000120
	NO 2000000372	A	20000321	NO	2000-372		20000125
	NO 318610	B1	20050418				
	US 6476035	B1	20021105	US	2000-491204		20000125
	BG 104148	A	20010531	BG	2000-104148		20000210
	BG 64904	B1	20060831				
	HK 1030220	A1	20041126	HK	2001-101274		20010221
	US 20030018050	A1	20030123	US	2002-223046		20020816
	US 6727263	B2	20040427				
	HK 1066806	A1	20070713	HK	2004-109852		20041213
	HK 1066807	A1	20070817	HK	2004-109853		20041213
PRI	ORITY APPLN. INFO.	:		DK	1997-892	A	19970725
				US	1997-53713P	P	19970725
				WO	1998-DK336	W	19980720

				US	2000-491204	A3	20000125

MARPAT 130:153571 OTHER SOURCE(S):

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The title compds. [I; X = O, S, CR4R5; Y = CR6R7, CR6R7CR8R9, CR6;CR7; XY = CR4;CR5, CR4;CR5CR6R7; Z = O, S; W = N, C, CH; A = II-IV; R1-R3,

R11-R17 = H, halo, CF3, etc.; R4-R9 = H, alky1; R11 = H, alky1, alkeny1, etc.]

their salts which are potent serotonin reuptake inhibitors and have 5-HTLR

5-HT1A

receptor antagonistic activity, were prepared Thus, treatment of
5-chioroindole with oxalyl chloride in Et2O followed by reaction of the
resulting 2-(5-chloro-1H-indol-3-yl)-2-oxoacetyl chloride with
1-(1,4-benzodioxan-5-yl)piperazine, and then reduction of the
intermediate

intermediate
with LIANH4 in THF afforded V.oxalate which showed IC50 of 5.0 nM against serotonin reuptake.

IT 220251-80-9P
RL RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of indole and 2,3-dihydroindole derivs. as potent serotonin

tonin
reuptake inhibitors and 5-HT1A receptor antagonists)
220251-80-9 CAPLUS
Ethanone, 2-(6-chloro-1H-indol-3-y1)-1-[4-(2,3-dihydro-1,4-benzodioxin-5-y1)-1-piperaziny1]- (CA INDEX NAME)

ANSWER 27 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

ANSWER 28 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 1998:402304 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 129:81760 129:16885a,16888a TITLE:

129:16885a,16888a
Preparation of spiro-substituted azacycles as
modulators of chemokine receptor activity
Mills, Sander G.; Springer, Martin S.; MacCoss,
Malcolm
Merck & Co., Inc., USA; Mills, Sander G.; Springer,
Martin S.; MacCoss, Malcolm
PCT Int. Appl., 297 pp.
CODEN: PIXXD2
Patent INVENTOR(S):

PATENT ASSIGNEE(S):

SOURCE

DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION;

	PAT	ENT I	NO.			KIN	D .	DATE			APPL	ICAT	ION	NO.		D	ATE	
							-									-		
	WO	98256	605			A1		1998	0618		WO 1	997-	US 23	586		1	9971	212
		W:	AL,	AM,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CU,	CZ,	EE,	GE,	GW,
			HU,	ID,	IL,	IS,	JP,	KG,	KR,	KZ,	LC,	LK,	LR,	LT,	LV,	MD,	MG,	MK,
			MN,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,
			US,	UZ,	VN,	YU												
		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,
			FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,	CF,	CG,	CI,	CM,
			GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG								
	AU	98580	033			A		1998	0703		AU 1	998-	5803	3		1	9971	212
OI	RITY	APPI	LN.	INFO	. :						US 1	996-	3273	5P		P 1	9961	213
											US 1	996-	3355	8P		P 1	9961	220
											GB 1	997-	3005			A 1	9970	213

WO 1997-US23586

W 19971212

(Continued)

MARPAT 129:81760 OTHER SOURCE(S):

L9 ANSWER 28 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Spiroazacycles I [R1 = H, alky1, aminoalky1, arylalky1, etc.; Q = \circ , S, S(\circ), S02, N; W = X bond, alky1, substituted alky1, etc.; YZ = fused

fused heteroaryl; m = n = 0 - 5 and m + n = 1 - 5] were prepared for use

as modulators of chemokine receptor activity (no data). Thus, spiroindoline II (R = 3,5-dimethylbenzoyl) was prepared starting from 3,5-dimethylbenzoic acid, 1,2-dihydro-1-(methylsulfonyl)spiro[3H-indole-3,4'-piperidine] monohydrochloride, and (S)-3,4-dichloro-N-methyl-β-2-propenylbenzeneethanamine.

II 167485-09-8P R.F. BAC (Biological activity or effector, except adverse); ESU (Riological)

RL: BAC (Biological activity or elector, and the file of the file

CN Ethanone,
1-[1,2-dihydro-1-(methylsulfonyl)spiro[3H-indole-3,4'-piperidin]1'-yl]-2-(5-fluoro-1H-indol-3-yl)- (CA INDEX NAME)

ANSWER 28 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

L9 ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1998:65894 CAPLUS
DOCUMENT NUMBER: 128:128015
ORIGINAL REFERENCE NO.: 128:25147a,25150a
TITLE: Preparation of amidinoindoles and amidinoazoles as inhibitors of factor Xa and of thrombin
INVENTOR(S): Dominguez, Celia; Han, Qi; Duffy, Daniel Emmett, Jeongsook Maria; Quan, Mimi Lifen; Rossi, Karen Wexler, Ruth Richmond
Du Pont Merck Pharmaceutical Co., USA
PCT Int. Appl., 176 pp.
CODEN: PIXXD2
Patent
English
1 PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. 3801428 A1 19980115 W0 1997-US11325 19970630
W: AM, AU, AZ, BR, BY, CA, CN, CZ, EE, HU, IL, VP, KG, KR, KZ, LT,
LV, MD, MX, NO, NZ, FL, RO, RU, SG, SI, SK, TJ, TM, UA, VN
RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT,

CA 2259573 A1 19980115 CA 1997-2259573 19970630 AU 9736456 A 19980202 AU 1997-36456 19970630 EP 960102 A1 19991201 EP 1997-93214 19970630 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE BUR 333696 A 20000623 NZ 1997-33596 19970630 AUTY AFPLN. INFO: US 1996-676766 A 19960708 PRIORITY APPLN. INFO.: US 1997-49519P P 19970613

WO 1997-US11325 W 19970630

MARPAT 128:128015 OTHER SOURCE(S):

The title compds. [I; W, W3 = CH, N; W1, W2 = C, CH, N (provided that or of W1 and W2 is C(C(=NH)NH2) and at most two of W, W1, W2, and W3 are N) one of D, Da = H, Cl-4 alkoxy, CN, etc. and the other is absent; one of

and Jb is substituted by -(CH2)n-Z-A-B; J, Ja, Jb combine to form an

DATE

ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) heterocyclic system contg. from 1-2 heteroatoms (N, O, and S), a heterocyclic ring wherein Jb = N and J and Ja = (un)substituted CH2, a heterocyclic ring wherein Jb = CH, J = (un)substituted NH and Ja = (un)substituted CH; Z = CH:CH, SOZCHZ, etc.; A = (un)substituted PhCH2, PhCH2CH2, etc.; B = CG-6 alkyl, (un)substituted PhCH2, 5-10 membered heterocyclic system, etc.], useful as inhibitors of factor Xa or mbin,

thrombin, were prepd. and formulated. Thus, reaction of 5-cyanoindole-1-acetic

were prepd. and formulated. Thus, reaction of 5-cyanoindole-1-acetic acid

acid

with 4-benzylpiperidine followed by treatment of the resulting

1-(4-benzylpiperidinocarbonyl)methyl-5-cyanoindole with RCl(g) in MeOH,

and then with (NR4)2CO3 in MeOH afforded the title compd. II. Some compds. I were evaluated and showed Ki of < 5 µM against thrombin.

IT 202123-90-8P 202123-94-2P 202123-96-4P

202123-90-8P 202123-94-2P 202124-28-5P

202124-04-7P 202124-24-1P 202124-28-5P

202126-86-1P

RL: BAC (Biological activity or effector, except adverse); BSU

(Biological

study, unclassified); SPN (Synthetic preparation); TRU (Therapeutic use);

BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amidinoindoles and amidinoazoles as inhibitors of factor Xa

and ef thrombin)

RN 202123-90-8 CAPLUS

CN 1B-Indole-5-carboximidamide, 3-[2-[4-[(4-methylphenyl)sulfonyl]-1-piperazinyl]-2-oxoethyl]- (CA INDEX NAME)

$$\mathsf{H}_2\mathsf{N}-\mathsf{C}$$

202123-94-2 CAPLUS 1H-Indole-5-carboximidamide, 3-[2-oxo-2-[4-[(phenylmethyl)sulfonyl]-1-phperazinyl]ethyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

202124-04-7 CAPLUS HA-Indole-5-carboximidamide, 3-[2-oxo-2-(4-phenyl-1-piperazinyl)ethyl]-(cA INDEX NAME)

202124-24-1 CAPLUS
IM-Indole-3-acetamide, 5-(aminoiminomethyl)-N-[2'-(aminosulfonyl)[1,1'-blphenyl]-4-yl]-N-methyl- (CA INDEX NAME)

$$\begin{array}{c} H \\ H_2N - C \\ NH \end{array}$$

202124-28-5 CAPLUS
1B-Indole-3-acetamide, 5-(aminoiminomethyl)-N-methyl-N-[2'[(methylamino)sulfonyl][1,1'-biphenyl]-4-yl)- (CA INDEX NAME)

202126-86-1 CAPLUS

ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN L9 (Continued)

202123-96-4 CAPLUS 1H-Indole-5-carboximidamide, 3-[2-oxo-2-[4-(phenylmethyl)-1piperazinyl]ethyl]- (CA INDEX NAME)

202123-97-5 CAPLUS
Glycine, N-[[5-(aminoiminomethyl)-1-methyl-1H-indol-3-yl]acetyl]-N-[[3(aminoiminomethyl)phenyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

202123-98-6 CAPLUS Glycine, N-[[5-(aminoiminomethyl)-1-methyl-1H-indol-3-yl]acetyl]-N-[[4-(aminoimnomethyl)phenyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Me} & \text{NH} \\ & \text{N} & \text{CH}_2 - \text{C-OMe} \\ & \text{H}_2 \text{N} - \text{C} \\ & \text{NH} \end{array}$$

202124-01-4 CAPLUS 1H-Indole-5-carboximidamide, 1-methyl-3-[2-oxo-2-[4-(phenylmethyl)-1-piperazinyl]ethyl]- (CA INDEX NAME)

ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) lH-Indole-5-carboximidamide, 3-[2-oxo-2-[4-(phenylmethyl)-1-piperidinyl]ethyl]- (CA INDEX NAME)

IT 202124-97-8
RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of amidinoindoles and amidinoazoles as inhibitors of factor Xa

factor Xa
and of thrombin)
RN 202124-97-8 CAPLUS
CN 1H-Indole-5-carbonitrile, 3-[2-oxo-2-(1-piperazinyl)ethyl]-,
hydrochloride
(1:1) (CA INDEX NAME)

● HCl

202124-91-2P Ri. RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of amidinoindoles and amidinoazoles as inhibitors of factor Xa

ox Na and of thrombin)
202124-91-2 CAPLUS
1HB-Indole-5-carbonitrile, 3-[2-[4-[(4-methylphenyl)sulfonyl]-1-piperazinyl]-2-oxoethyl]- (CA INDEX NAME)

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

A 19960224

A 19960904

W 19970220

A3 19971021

A3 19990730

L9 ANSWER 29 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) L9 ANSWER 30 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1997:579718 CAPLUS DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 127:248104 127:48481a TITLE: Preparation of aryloxooxazolidinylmethylacetamides related compounds as antibacterials. Gravestock, Michael Barry Zeneca Ltd., UK; Gravestock, Michael Barry PCT Int. Appl., 111 pp. CODEN: PIXXD2 Patent INVENTOR (S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE. DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English PATENT NO. KIND DATE APPLICATION NO. W0 9730995 A1 19970828 W0 1997-GB462 19970220
W1 AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, CE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MN, NN, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG

ZA 9701699 A 19970925 ZA 1997-1469 19970220
AU 9718053 A 19970910 AU 1997-18053 19970220
EP 882042 A1 19981209 EP 1997-903509 19970220
R: CB, DE, FR, BJ, LT, LT
JF 11514662 T 19991214 JP 1997-52988 19970220
N 1997060443 A 2005311 IN 1997-B443 19970220 JP 1997-529888 IN 1997-DE443 US 1997-945160 US 1999-364389 US 2001-836095 IN 1997DE00443 US 5981528 US 6271383 19970221 19971021 19990730 20050311 19991109

20010807 20020402

MARPAT 127:248104

GB 1996-3939

GB 1996-18404

WO 1997-GB462

US 1997-945160

US 1999-364389

Bl

PRIORITY APPLN. INFO.:

OTHER SOURCE (S):

L9 ANSWER 30 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Title compds. (I; R1 = OH, C1, Br, F, alkylsulfonyloxy, amino, N3,

prepared
Thus, a mixture of tert-Bu 1,2,3,6-tetrahydro-4(trifluoromethylsulfonyloxy)pyridine-1-carboxylate,
Pd2(dibenzylideneacetone)2, Ph3As, and LiCl in N-methylpyrrolidine was
treated with (8)-5-acetamidomethyl-3-(4-trimethyltinphenyl)oxazolidin-2one (preparation given) followed by stirring at room temperature to 40°

give 23% (S)-N-[3-[4-(1-tert-butyloxycarbonyl-1,2,5,6-tetrahydropyrid-4-yl)phenyl]-2-oxooxazolidin-5-ylmethyl]acetamide. The latter showed a min.

inhibitory concentration of 1.0 µg/mL against Staphylococcus aureus Oxford. IT 195816-92-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

ogical
study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of aryloxocxazolidinylmethylacetamides and related compds. as as antibacterials)

antibacterials/ 19816-92-3 CAPUS Acetanide, N-[(i5s)-3-[4-[1-[2-(5-fluoro-1H-indol-3-yl)acetyl]-1,2,3,6-tetrahydro-4-pyridinyl]phenyl]-2-oxo-5-oxazolidinyl]methyl]- (CA INDEX

Absolute stereochemistry.

ANSWER 30 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 31 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 1997:456960 CAPLUS MENT NUMBER: 127:95194 INAL REFERENCE NO.: 127:16329a,18332a ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

TITLE:

127:18329a,18332a

New benzisoindole derivatives as inhibitors of farnesyl transferase, their preparation, and pharmaceutical compositions containing them.

Commercon, Alain; Lebrum, Alain; Mailliet, Patrick; Feyronel, Jean Francois; Sounigo, Fabienne; Truchon, Alain; Zucco, Martine; Cheve, Michel Rhone-Poulenc Rorer SA, Fr.

Fr. Demande, 36 pp.

CODEN: FRXXBL
Patent
French
1 INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION:

		TENT I																	
	FR	2736	541			A1		1997	0117		FR	19	95-8	8296			1	9950	710
	FR	2736	541			Bl		1997	0822										
	TW	2736 4387 1996	92			В		2001	0607		TW	19	96-8	8510	8158		1	9960	705
	IN	1996	DE01	492		A		2005	0311		IN	19	96-1	DE14:	92		1	9960	705
	CA	2224	414			A1		1997	0130		CA	19	996-2	2224	414		1	9960	708
	WO	97031	050			A1		1997	0130		WO	19	996-1	FR10	62		1	9960	708
		W:						CA,											
								MK,											
								AM,											
		RW:																	
								PT,	SE,	BF,	B	,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,
			MR,	NE,	SN,	TD,	TG												
	AU	9665: 7121:	224			A		1997	0210		AU	19	996-6	6522	4		1	9960	708
	AU	7121	94			B2		1999	1028										
		8391									EP	19	996-9	9249.	52		1	9960	708
		8391																	
		R:	,			,		,					,	,					
FI	-027	1100	700									2.0		0054				00.50	700
	CIN	1190	440			A		T330	1012		CIV	13	220	1954.	13		Т	9900	100
	CIN	1096	110					2002.	1510			20	006	E 0 E E	c 7		2	0050	700
	3.00	1057	1123			1		1000	1015		3.00	13	770-	2022	57		1	2200	700
	D.C	2120	272			4.2		2000	0201		TO.	10	006	2242	52		1	2250	700
	TI.	1228	10			2		2000	0701		TI.	10	996-	1008	12		1	9960 9960	700
	CV	1190: 1096: 1151: 1853: 2139: 1228: 2822: 2916: 9605: 9609:	50			D.C.		2001	1202		CK	10	100	25	12		1	9950	70.9
	CZ	2916	20			B6		2001	0416		CZ	10	998_	54			1	9960	708
	73	9605	868			2		1997	0129		73	10	996-	5868			1	9960	710
	BR	9609	440			A		19991	0629		BR	19	996-9	344N			1	9960	710
	NO	98001	194			Δ.		1998	0023		NIO.	10	998_9	94			1	9980	109
	NO	9800 3095	55			B1		2001	0219		140						_	,,,,,	100
	US	59361	397			A		19991	0810		US	19	998-9	9818	40		1	9980	723
	GR	3031	409			Т3		2000	0131		GR	19	999-	4020	01		1	9991	007
PRIC	RITS	APP:	INI.	TNFO							FR	19	95-1	8296			A 1	9950	710
																			•

ANSWER 31 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN methyl ester, (3aR,4S,9S,9aR)-rel- (CA INDEX NAME) (Continued)

Relative stereochemistry.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

logical
study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of new benzisoindole derivs. farnesyl transferase

(preparation of new benzisoindole derivs. farnesyl transferase inhibitors)
RN 191989-23-8 CAPLUS
CN 4,9-Ethano-3aH-benz[f]isoindole-3a-carboxylic acid,
2-[2-[5-brono-1H-indol-3-y1)acety1]-1,2,3,4,9,9a-hexahydro-9-phenyl-,
(3aR,4S,9S,9aR)-rel- (CA INDEX NAME)

Relative stereochemistry,

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT: FORMAT

L9 ANSWER 31 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN OTHER SOURCE(S): MARPAT 127:95194 (Continued)

Title compds. I [R = (un)substituted (CH2)mX1(CH2)nZ; X1 = bond, O, S; m

= 0-1; n = 0-2; Z = CO2H, alkoxycarbonyl, (un)substituted carbamoyl, etc.; R1, R2 = H, halo, alkyl, (un)substituted alkoxy; or R1R2 form (un)saturated

(un)saturated

(un)setricted in the strength of the st

claimed. The compds. are inhibitors of farnesyl transferase, and show marked antitumor and antileukemic properties. For example, cis-3,6-diphenyl-1,4-cyclohexadienecarboxylic acid Me ester (preparation

n) reacted with FhCH2N(CH2OBu)(CH2SiMe3) in refluxing CF3CO2H to give the intermediate hexahydroisoindole derivative II.HCl, which was further cvclized

used by CF3SO3H at 5-20° to give the benz[f]isoindole intermediate III. This was then converted in 3 steps to title compound IV. In an assay for inhibition of farnesyl transferase, IV had an IC50 of 0.31 µM. 191989-96-5P

191999-96-5F RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate; preparation of new benzisoindole derivs. farnesyl

transferase

inhibitors)

inhibitors;
191989-96-5 CAPLUS
4,9-Ethano-3aH-benz[f]isoindole-3a-carboxylic acid,
2-[2-(5-bromo-1H-indol-3-yl)acetyl]-1,2,3,4,9,9a-hexahydro-9-phenyl-,

L9 ANSWER 32 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1995:995279 CAPLUS
DOCUMENT NUMBER: 124:145907
CRIGINAL REFERENCE NO.: 124:27133a, 27136a
TITLE: Preparation of 1-(3-indolylalkyl)-4-(3-indolyl)piperidines as dopamine agonists or antagonists.

INVENTOR(S): Boetcher, Henning, Maerz, Joachim, Seyfried, Christoph; Greiner, Hartmut; Bartoszyk, Gerd Merck Patent Gabh, Germany
CODE: CODE: GRWXBX
DOCUMENT TYPE: CODE: GRWXBX
FAMILY ACC. NUM. COUNT: 1
FAMILY ACC. NUM. COUNT: 1
FATENT INFORMATION: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.			PLICATION NO.	
DE 4414113			1994-4414113	
EP 683166	A1 199		1995-105227	
EP 683166	B1 199	81028		
		FR. GB. G	R, IE, IT, LI, LU,	NL. PT. SE
AT 172730 ES 2125508 AU 9516488	T 199		1995-105227	
ES 2125508	T3 199	90301 ES	1995-105227	19950407
AU 9516488	A 199	51102 AU	1995-16488	19950413
AU 697749	B2 199	81015		
JP 07291969	A 199	51107 JP	1995-91077	19950417
SK 280881	B6 200	000814 SK	1995-508 1995-2147451	19950419
CA 2147451	A1 199	51023 CA	1995-2147451	19950420
CA 2147451	C 200	60328		
CN 1114651	A 199	60110 CN	1995-104705	19950420
CN 1047385	C 199 B 200	91215		
			1995-84103916	
NO 9501529	A 199	51023 NO	1995-1529	19950421
NO 307831	B1 200	000605		
ZA 9503260	A 199	60109 ZA	1995-3260	19950421
HU 74096			1995-1139	
US 5693655	A 199	71202 US	1995-426405	19950421
CZ 285369	B6 199		1995-1035	
RU 2151148		000620 RU	1995-106675	19950421
PL 180781	B1 200	10430 PL	1995-308287	19950421
PRIORITY APPLN. INFO.:		DE	1994-4414113	A 19940422
OTHER SOURCE(S):	CASREACT 1	24:145907; 1	MARPAT 124:145907	
OT.				

L9 ANSWER 33 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1995:851691 CAPLUS

123:285765 123:51207a, 51210a

ANSWER 32 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
Title compds. [I; R1-R4 = H, alkyl, OH, alkoxy, F, Cl, Br, iodo, cyano, CF3, CO2H, CONH2, alkoxycarbonyl, etc.; R1R2, R3R4 = OCH2O; R5 = H, OH;

= H; R5R6 = bond; n = 2-6], were prepared as drugs (no data). Thus, 3-(4-chlorobutyl)-5-methoxyindole and 4-(3-indolyl)piperidine were refluxed 8 h in MeCN to give 3-[1-[4-(5-methoxyindol-3-yl)butyl]-4-piperidinyl]indole hydrochloride. 173150-68-0 173150-69-1 RI.; RCT (Reactant); RACT (Reactant) regretation of 1-(3-indolylalkyl)-4-(3-indolyl)piperidines as nine.

name agonists or antagonists)
173150-68-0 CAPLUS
Ethanone, 2-(5-fluoro-1H-indol-3-yl)-1-[4-(5-fluoro-1H-indol-3-yl)-1piperidinyl]- (CA INDEX NAME)

173150-69-1 CAPLUS Ethanone, 2-(5-fiuoro-1H-indo1-3-y1)-1-[4-(4-fiuoro-1H-indo1-3-y1)-1-piperidiny1]- (CA INDEX NAME)

ANSWER 33 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) (prepn. of perhydroisoindole antiemetics) 153438-63-2 CAPLUS (Proposition of the continued) 153438-63-2 CAPLUS (Proposition of the continued of the c

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

123:120/A,51210a
Preparation of perhydroisolndole antiemetics
Garret, Claude; Louvel, Erik
Rhone-Poulenc Rorer S.A., Fr.
PCT Int. Appl., 62 pp.
CODEN: PIXXD2 PATENT ASSIGNEE(S): DOCUMENT TYPE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: MO 9509628 A1 19950413 WO 1994-FR1160 19941005
W: AM, AU, BB, BG, BR, BY, CA, CM, CZ, EE, FI, GE, HU, JP, KG, KP,
KK, KZ, LK, LR, LT, LV, MD, MG, MN, NO, NZ, FL, RO, RU, SI, SK,
TJ, TT, UA, US, UZ, VN
RW: KE, MW, SD, SZ, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU,
MC, NL, FT, SE, BF, BJ, CF, CC, CI, CM, GA, GN, ML, ME, NE, SN,
TD, TG
FR 2710842 A1 19950414 FR 1993-11945 19931007
FR 2710842 B1 19951124
AU 9478581 A 19950501 AU 1994-78581
PRIORITY APPLN. INFO:

OTHER SOURCE(S): CASREACT 123:285765; MARPAT 123:285765

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

TITLE: INVENTOR(S):

The title compds. [I; R = (un)substituted Ph; RI = (un)substituted Ph; cyclohexadienyl, naphthyl, indenyl, (un)substituted heterocyclyl; R2 = H; halogen, OH, alkyl, aminoalkyl, alkylaminoalkyl, dialkylaminoalkyl, alkylaminoalkyl, alkylaminoalkyl, alkylaminoalkyl, benzyloxycarbonyl, NH2, acylamino; R3 = (un)substituted alkyloxycarbonyl, benzyloxycarbonyl, NH2, acylamino; R3 = (un)substituted Ph; R4 = OH or F if R5 = H; etc.] [e.g., (3a8, 48, 7a8)-7, 7-diphenyl-4-(2-methoxyphenyl)-2-tert-butoxycarbonyl-4-perhydroisoindololl, useful as antiemetics, are prepared and I-containing formulations presented.

1334,35-63-47 RE: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

L9 ANSWER 34 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1995:781772 CAPLUS
DOCUMENT NUMBER: 123:169671
ORIGINAL REFERENCE NO.: 123:30303a,30306a
TITLE: Preparation of spirocyclic compounds as neurokinin antagonists
INVENTOR(S): MacCoss, Malcolm; Mills, Sander G.; Shah, Shrenik K.; Chiang, Yuan-Ching P.; Dunn, Patrick T.; Koyama, Hiros; Finke, Paul E.; Qi, Hongbo; Robichaud, Albert J.

PATENT ASSIGNEE(S): SOURCE:

J.
Merck and Co., Inc., USA
PCT Int. Appl., 226 pp.
CODEN: PIXXD2
Patent
English 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	ENT 1	NO.			KINI	D :	DATE			APF	LIC	CAT:	ION :	NO.		D	ATE	
							-										_		
	WO	9429	309			A1		1994	1222		WO	199	94-1	JS55	45		1	9940	517
		W:	AU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	FI	, 1	ΗU,	JP,	KR,	KZ,	LK,	LV,	MG,
			MN,	MW,	NO,	NZ,	PL,	RO,	RU,	SD,	SI	, :	SK,	TT,	UA,	US,	UZ		
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	١, ١	IE,	IT,	LU,	MC,	NL,	PT,	SE,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	MI	., 1	MR,	NE,	SN,	TD,	TG		
	CA	2163	995			A1		1994	1222		CA	199	94-	2163	995		1	9940	517
	ΑU	9472	011			A		1995	0103		ΑU	199	94-	7201	1		1	9940	517
	ΑU	6800	20			B2		1997	0717										
	EP	7026	81			A1		1996	0327		EΡ	199	95-5	9019	79		1	9940	517
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	٠, :	ΙE,	IT,	LI,	LU,	NL,	PT,	SE
	JP	0851	1522			T		1996	1203		JP	199	94-	5018	02		1	9940	517
	ZA	9403	946			A		1995	0120		ZA	199	94-	3946			1	9940	606
PRIOF	(TI	APP:	LN.	INFO	. :						US	199	93-	7290	4		A 1	9930	607
											wo	199	94-1	JS 5 5	45	,	w 1	9940	517

MARPAT 123:169671 OTHER SOURCE(S):

L9 ANSWER 34 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Spirocyclic nitrogen-heterocyclic compds. were disclosed as tachykinin receptor antagonists useful for the treatment of inflammatory diseases, pain or migraine, and asthma. In particular, said compds. were shown to be neurokinin antagonists. Many example compds. are claimed. One such specific compound is N-[3-(3,4-dichlorophenyl)-4-[1,2-dihydro-1-(sulfonylmethyl)spiro[3H-indole-3,4'-piperidin]-1'-yl]butyl]-2,2-dimethylpropanamide (I).
167485-09-8P (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); FREF (Preparation); USES (Uses) (preparation of spirocyclic compds. as kinin receptor antagonists) 167485-09-8 CAPLUS Ethanone,

IN 10/10/00/00 CALIBOO (C) Ethanone, 1-(1,2-dihydro), 1-(1,2-dihydro), 1-(methylsulfonyl), spiro[3H-indole-3,4'-piperidin]-1'-yl]-2-(5-fluoro-1H-indol-3-yl)- (CA INDEX NAME)

ANSWER 34 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

REFERENCE COUNT:

L9 ANSWER 35 OF 44 CAPLUS COPYRIGHT 2009 ACS ON STN
ACCESSION NUMBER: 1995:772570 CAPLUS
DOCUMENT NUMBER: 123:169499
123:169499
1171LE: 123:0255a, 30255a
1171LE: 1ndole derivatives as 5-HT1-like agonists for use in migraine
Wythes, Martin James
PATENT ASSIGNEE(S): Pfizer Ltd., UK, Pfizer Inc., Pfizer Research and Development Company, N.V./S.A.
SOURCE: CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1 DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	TENT NO																DATE	
	942412																	
	W: 3	AU, B	R,	CA,	CN,	CZ,	FI,	HU,	JP,	K	١,	NO,	NZ,	PL,	RU,	US	3	
	RW: A	AT, B	E,	CH,	DE,	DK,	ES,	FR,	GB,	GI	١,	IE,	IT,	LU,	MC,	N	, PT,	SE
CA	215739	9.7			A1		1994	1027		CA	19	94-	2157	397			19940	0411
CA	215739 946563	7			C		1999	0706										
AU	946567	70			A		1994	1108		ΑU	19	94-	6567	0			19940	0411
	940648																	
EP	695301				A1		1996	0207		EP	19	94-	9135	73			19940	0411
EP	695301	1			B1		1996	1030										
	R: 1	AT, B	E,	CH,	DE,	DK,	ES,	FR,	GB,	GI	١,	IE,	IT,	LI,	LU,	N	, PT,	SE
CN	112134 085070	18			A		1996	0424		CN	19	94-	1918	50			19940	0411
JP	085070	183			T		1996	0730		JP	19	94-	5227	26			19940	0411
HU	73807				A2		1996	0930		HU	19	95-	1920				19940	111
AT	144773	3			T		1996	1115		AT	19	94-	9135	73			19940	0411
ES	209465	3			Т3		1997	0116		ES	19	94-	9135	73			19940	0411
ZA	940272	22			A		1995	1020		ZA	19	94-	2722				19940	1420
FI	950494	14			A		1995	1017		FI	19	95-	4944				19951	1017
NO	950416																19951	
US	560798	0			A		1997	0304		US	19	95-	5325	73			19951	1020
PRIORIT	Y APPLE	J. IN	FO.	:						GB	19	93-	9360			Α	19930	1422
										GB	19	93-	2443	3		Α	19931	1127
																	7004	
										WO	Т3	94-	RETT	21		W	19940	J411

OTHER SOURCE(S): MARPAT 123:169499

The title compds., 3-(pyrrolidinylmethyl)indoles and 3-(piperidinylmethyl)indoles I [Rl = (2-pyrrolidinyl)methyl,

L9 ANSWER 35 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

3-pyrrolidiny1, 4-piperidiny1, (3-piperidiny1)methy1; R2 = alky1,
oxoalky1, etc.] were disclosed as selective 5-HT1-11ke agonists useful in
the treatment of migraine, cluster headache, chronic paroxysmal
hemicrania
and headache assocd. with vascular disorders. A specifically claimed
example compd. is 5-(3-hydroxybuty1)-3-[(R)-(1-methy1-2pyrrolidiny1)methy1]-1-H-indole (II).

IT 167303-72-2P
RL! RCT (Reactant); SRN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(preparation of (aminoalky1)indoles 5-HT1-like agonists)
RN 167303-72-2 CAPLUS
CN 1H-Indole-3-acetamide, 5-bromo-N-methy1-N-(phenylmethy1)- (CA INDEX
NAME)

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE 3 REFERENCE COUNT: FORMAT

L9 ANSWER 36 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1995:615038 CAPLUS DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 123:32956 123:6087a,6090a

123:6087a,6090a
Preparation of pharmaceutical perhydroisoindole
derivatives as neurokinin A antagonists
Crespo, Andre; Fardin, Veronique; Guillaume,
Jean-Marc; Malleron, Jean -Luc; Peyronel,
Jean-Francois
Rhone-Poulenc Rorer S.A., Fr.
PCT Int. Appl., 43 pp.
CODEN: PIXXD2
Patent
French
1 TITLE: INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION;

PA:	TENT NO	٠.			KIN)	DATE			API	PLIC	CAT	ION	NO.		Ε	ATE	
WO	942282	22			A1		1994	1013		WO	199	94-	FR37	1		1	9940	401
	W: A	λU,	CA,	CZ,	FI,	HU,	JP,	KR,	NO,	N2	Ζ, Ι	PL,	RU,	SK,	UA,	US		
	RW: A	AΤ,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GI	R, :	IE,	IT,	LU,	MC,	NL,	PT,	SE
FR	270367	79			A1		1994	1014		FR	199	93-	3965			1	9930	405
FR	270367	79			B1		1995	0623										
CA	215866	53			A1		1994	1013		CA	199	94-	2158	663		1	9940	401
AU	946506	8			A		1994	1024		ΑU	199	94-	6506	8		1	9940	401
EP	693059	9			A1		1996	0124		EP	199	94-	9125	82		1	9940	401
EP	693059	9			B1		1997	0312										
	R: 7	AΤ,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GE	R, 3	IE,	IT,	LI,	LU,	NL,	PT,	SE
JP	085082	283			T		1996	0903		JP	199	94-	5217	62		1	9940	401
HU	74089				A2		1996	1128		ΗU	199	95-	2902			1	9940	401
AT	150014	2			T		1997	0315		AT	199	94-	9125	82		1	9940	401
ES	209960	1			Т3		1997	0516		ES	199	94-	9125	82		1	9940	401
US	563127	79			A		1997	0520		US	199	95-	4484	02		1	9950	607
NO	950391	13			A		1995	1002		NO	199	95-	3913			1	9951	002
FI	950473	30			A		1995	1117		FI	199	95-	4730			1	9951	004
PRIORITY	APPLN	۷.	INFO							FR	199	93-	3965			A 1	9930	405
										WO	199	94-1	FR37	1		W 1	9940	401

OTHER SOURCE(S): MARPAT 123:32956

ANSWER 36 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT: FORMAT

ANSWER 36 OF 44 CAPLUS COFYRIGHT 2009 ACS on STN (Continued)
Title compds, I (R = (substituted)Ph; Rl = (substituted)Ph, PhCh2O,
(substituted)-Cl-4 alkyl, (substituted)amino, (substituted)heterocyclyl,
cyclohexadienyl, naphthyl, indenyl; R2 = H, halo, HO, alkyl, aminoalkyl,
allylaminoalkyl, dlalkylaminoalkyl, etc.; R3 = (substituted)Ph), are
prepared (3AR,4R,5R,7AB)-7,7-diphenyl-4-(2-methoxyphenyl)perhydro-4,5isoindolediol (preparation given) and 3-indolylacetic acid in CH2Cl2
added

added
to 1-benzotriazolylol hydrate, 1-(3-dimethylaminopropyl)-3ethylcarbodiimide and diisopropylethylamine to give (3aR,4R,5E,7aR)-I (R1
= 3-indolyl, R2 = H, R3 - 2-(MeO)CEH4) which at 10-1000 mM on human
receptor NK2 showed IC50 of 215 nM. A formulation tablet comprising I is
given.
153838-54-8P 163838-57-IP

IT 163838-54-8P 163838-57-1P
RL: RAC (Biological activity or effector, except adverse); BSU
(Biological study); SPN (Synthetic preparation); THU (Therapeutic use);
BTOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of pharmaceutical perhydrolsoindole derivs. as neurokinin λ
antagonists)
RN 163838-54-8 CAPLUS
CN 1H-Isoindole-4,5-diol,
octahydro-2-[(5-hydroxy-1H-indol-3-y1)acety1]-4-(2-methoxyphenyl)-7,7-dlphenyl-, [3aR-(3aα,4β,5β,7aα)](9CI) (CA INDEX NAME)

163838-57-1 CAPLUS
Ethanone, 2-(5-fluoro-1H-indo1-3-y1)-1-[(3aR, 4R, 5R, 7aR)-octahydro-4,5-dihydroxy-4-(2-methoxypheny1)-7,7-dipheny1-2H-isoindo1-2-y1]-, rel-

Relative stereochemistry.

L9 ANSWER 37 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1994:270102 CAPLUS
DOCUMENT NUMBER: 120:270102
CRIGINAL REFERENCE NO: 120:47843a, 47846a
TITLE: antaqonists and their preparation
Achard, Danlel; Grisoni, Serge; Malleron, Jean Luc;
PATENT ASSIGNEE(S): Rhome-Poulenc Rozer S.A., Fr.
SOURCE: CODEN: FIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 1
FATENT INFORMATION:

PATENT INFORMATION:			
PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9321155	A1 19931028	WO 1993-FR352 KZ, NO, NZ, PL, RU, SK,	19930408
RW: AT, BE, C	H, DE, DK, ES, FR,	GB, GR, IE, IT, LU, MC,	NL, PT, SE
FR 2689888	A1 19931015	FR 1992-4390	19920410
FD 2689888	B1 19940610		
IL 105255	A 19970218	IL 1993-105255	19930401
ZA 9302527	A 19931108	ZA 1993-2527	19930408
AU 9339565	A 19931118	IL 1993-105255 ZA 1993-2527 AU 1993-39565	19930408
AU 667214	B2 19960314		
EP 635003	A1 19950125	EP 1993-909005	19930408
EP 635003	B1 19980617		
R: AT, BE, C	, DE, DK, ES, FR,	GB, GR, IE, IT, LI, LU,	NL, PT, SE
JP 07505410	T 19950615	JP 1993-518041	19930408
JP 3205557	B2 20010904		
HU 71354	A2 19951128	HU 1994-2911	19930408
PL 172754	B1 19971128	PL 1993-305360	19930408
SK 279032	B6 19980506	SK 1994-1220 AT 1993-909005	19930408
AT 167472	T 19980715	AT 1993-909005	19930408
CZ 284213	B6 1998 0 916	CZ 1994-2482	19930408
ES 2118232	T3 19980916	ES 1993-909005	19930408
RU 2127260	C1 19990310	RU 1994-45855 NO 1994-3692	19930408
NO 9403692	A 19941003	NO 1994-3692	19941003
FI 9404729	A 19941007	FI 1994-4729	19941007
FI 105023	Bl 20000531		
US 5484804	A 19960116	US 1994-313121 FR 1992-4390	19941011
PRIORITY APPLN. INFO.:		FR 1992-4390	19920410
		WO 1993-FR352	19930408
OTHER COMPCE(C).	MARDAT 100,0001	02	

MARPAT 120:270102

L9 ANSWER 37 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Title compds, I [R = Ph optionally substituted with halogen or Me in position 2 or 3; R1 = (un)substituted Ph, cyclohexadienyl, naphthyl, indenyl, heterocyclyl; R2 = H, halo, OH, alkyl, aminoalkyl, CO2H, amino, etc.; R3 = Ph optionally substituted in position 2 by C1-2 alkyl or alkoxy; R4 = F, OH; E5 = H; or R4 = R5 = OH; or R4R5 = bond| and their stereoisomers, isomer mixts., and salts, are claimed (40 synthetic examples). For example, N-acylation of [3a(S), 4(S), 7a(S)]-7, 7-diphenyl-4-(2-methoxyphenyl)perhydroisoindol-4-ol (prepared in 4 steps) with (S)-2-(MeO)C6H4CHMeCO2H (prepared in 3 steps) of

(prepared in 4 steps, with the first part of the

guinea pigs. II 153438-63-2P RL: BAC (Biological activity or effector, except adverse); BSU (Biological

logical study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as substance P antagonist) 154438-63-2 CAPLUS 154438-63-2 CAPLUS 18-Isoindol-4-ol, 2-[(5-fluoro-1H-indol-3-y1)acety1]octahydro-4-(2-methoxyphenyl)-7,7-diphenyl-, [3aS-(3aα,4β,7aα)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 37 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

L9 ANSWER 38 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1994:244664 CAPLUS
DOCUMENT NUMBER: 120:244664
ORIGINAL REFERENCE NO.: 120:34361a, 43364a
TITLE: PATENT ASSIGNEE (S): Achard, Daniel; Grisoni, Serge; Malleron, Jean Luc; Peyronei, Jean Francois; Tabart, Michel
Rhone-Poulenc Rorer S.A., Fr.
SOURCE: CODEN: PIXXD2
DOCUMENT TYPE: PATENT ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

						D DATE						
						19931028						
						HU, JP, KR,						040
						DK, ES, FR,						. s
TD	26898	99			8.1	19931016	_ TD					
FR	26898	89			B1	19940610						
IL	10525	6			A	19970814	IL	1997-1	05256		19930	040
ZA	93025	28			A	19931028	ZA	1993-2	528		19930	040
						19931118						
						19960321						
EP	63500	12			A1	19950125	EP	1993-9	09004		19930	040
EP	63500	12			B1	19980722						
	R:	AT,	BE,	CH,	DE,	DK, ES, FR,	GB, G	R, IE,	IT, LI,	LU,	NL, PT,	, 5
JP	07505	409			T	19950615	JP	1993-5	18040		19930	040
HU	71330				A2	19951128	HU	1994-2	912		19930	040
						19971128						
AT	16867					19980815						
	21189					19981001						
RU	21204	138			C1	19981020	RU	1994-4	5867		19930	040
CZ	28459					19990113						
	94037					19941005						
						19941007		1994-4	728		1994	100
						20000531						
						19951031						
DRITY	APPI	N.	INFO.	. :			FR	1992-4	391	I	19920	041
							WO	1993-F	R351	7	19931	าสก

OTHER SOURCE(S): MARPAT 120:244664

L9 ANSWER 38 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Title compds. (I; R = Ph, 2- or 3-halophenyl, -methylphenyl; R1 = Ph, 2-methyl- or -ethylphenyl, -methoxy- or -ethoxyphenyl; R2 = F, OH; R3 = $\frac{1}{2}$

OH; R2R3 = bond; R4 = H, protective group) were prepared. Thus, (3aRS, 7aRS) - 7, 7-diphenylperhydroisoindol-4-one was converted in 3 steps

(3aRS,7aRS)-7,7-diphenylperhydrolsoindol-4-one was converted in 3 steps to (S,S)-I (R = Ph, R1R2 = O, R3 = H, R4 = CO2CMe3) which was condensed with the Grignard reagent from 2-(MeO)C6H4Br to give, after deprotection, isoindolol II (R4 = H). The latter was condensed with (S)-2-(MeO)C6H4CMECO2H (preparation given) to give II (R4 = S)-2-(MeO)C6H4CMECO2H (preparation) to give II (R4 = S)-2-(MeO)C6H4CMECO2H (preparation) ISSI (S)-2-(MeO)C6H4CMECO2H (preparation) (BSS (Uses)) (Biological study); PREP (Preparation); USES (Uses) (preparation of, as substance P antagonist)
RN 13345-63-2 CAPLUS

Absolute stereochemistry.

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

ANSWER 39 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 1993:671015 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 119:271015 119:48497a,48500a 119:48497a,48500a (Indolylethyl)piperidine NK2 receptor antagonists Cooper, Anthony William James; Hagan, Russell Michael Glazo Group Ltd., UK PCT Int. Appl., 39 pp. CODEN: FIXXD2 Patent TITLE: PATENT ASSIGNEE(S): DOCUMENT TYPE: English FAMILY ACC, NUM. COUNT: PATENT INFORMATION: PATENT NO. A2 19930722 W0 1993-EP1U1
A3 19931014
A3 19931014
CG, CI, CM, CA, CN, ML, MR, SN, TD, TG
A 19930803 AU 1993-3310
GB 1992-1179

V0101 KIND DATE APPLICATION NO. DATE WO 9314084
WO 9314084
RW: AT, BE, CH,
BF, BJ, CF,
AU 9333513
PRIORITY APPLN. INFO.: 19930115 NL, PT, SE, 19930115 19920121 A 19930115

The title compds. I [R1 = (un)substituted Ph; R2 = H, H0, C1-4 alkoxy; R3 = H, C1-4 alkyl; R4 = H, C1-4 alkyl; C1-4 alkoxy; R5 = H, C1-4 alkyl; AB

CN, halogen, n=0-2], useful in the treatment of conditions mediated by tachykinins, including NKA, NKB, and substance P, acting at the NK2 receptor, are prepared Thus, (R)-methylphenyl sulfoxide was reacted 1.1 with Li

Li bis (trimethylsilyl)amide, and the intermediate reacted with 1-[5-fluoro-1H-indol-3-yl)ethyl]-4-piperidone, followed by

MARPAT 119:271015

unesulfonic acid, producing (R)-1-[2-(5-fluoro-1H-indol-3-y1)ethy1]-4-[(phenylsulfiny1)methy1]-4-piperidinol methanesulfonic acid salt (II).

ΙI demonstrated anxiolytic activity in the mouse light-dark box and the rat elevated plus-maze. 151191-69-4P 151191-70-7P 151191-71-8P

ANSWER 39 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) Ethanone, 2-(5-fluoro-lH-indol-3-yl)-1-[4-hydroxy-4-[[(2-methylphenyl)thio]methyl]-1-piperidinyl]- (CA INDEX NAME)

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 39 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
151191-75-2P 151191-78-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREF (Preparation); RACT
(Reactant or reagent) (prepn. and reaction of, in prepn. of NK2 receptor antagonists) 151191-69-4 CAPLUS eridinone, 1-[2-(5-fluoro-1H-indol-3-yl)acetyl]- (CA INDEX NAME)

151191-70-7 CAPLUS Ethanone, 2-(5-fluoro-1H-indol-3-y1)-1-[4-hydroxy-4-[(phenylsulfiny1)methy1]-1-piperidiny1]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ &$$

151191-71-8 CAPLUS Ethanone, 2-(5-fluoro-1H-indol-3-yl)-1-[4-hydroxy-4-[[(2-methylphenyl)sulfinyl]methyl]-1-piperidinyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

151191-75-2 CAPLUS Ethanone, 2-(5-fluoro-1H-indol-3-yl)-1-[4-hydroxy-4-{{(2-methylphenyl)sulfonyl]methyl}-1-piperldinyl]- (CA INDEX NAME)

151191-78-5 CAPLUS

ANSWER 40 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN SSION NUMBER: 1993:168924 CAPLUS PLUS COPYRIGHT 2009 ACS on STN
1993:168924 CAPLUS
118:168924 TIB:28969a, 28972a
Search for B-adrenoblockers among aminooxypropyl
derivatives of 4-hydroxyindolylacetic acid and
4-hydroxyskatole
Glushkov, R. G.; Mashkovskii, M. D.; Skryabin, G. K.;
Suvorov, N. N., Kozlovskii, A. G.; Vinograd, L. Kh.;
Yurhakov, S. D.; Arinbasarov, M. U.; Tribunskaya, Yu.
I.; et al.
TSKhLS, VMIKhFI im. S. Ordzhonikidze, Moscow, Russia
Khimiko-Farmatsevticheskii Zhurnal (1992), 26(6),
18-21 ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.: AUTHOR(S): CORPORATE SOURCE: 18-21 CODEN: KHFZAN; ISSN: 0023-1134 Journal Russian CASREACT 118:168924

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

AB Treating indoles I (R = CH2CO2Me, Me, CH2CONH2, CH2CONMe2) with 2-(chloromethyl)oxirane gave 74-82.5% glycidyloxy derivs. which were substituted by Me2CHN12 and Me3CNH2 to give 60.5-94.5% aminohydroxypropoxy derivs. II (R1 = Me2CH, CMe3). The highest blocking activity was displayed by II (R = Me, R1 = CMe3) and by II (R = CH2CO2Me, R1 = CMe3). IT 145101-52-6

145101-32-6
RL: PROC (Process)
(substitution of, by epichlorohydrin)
145101-52-6 CAPUS
1H-1ndole-3-acetamide, 4-hydroxy-N,N-dimethyl- (CA INDEX NAME)

10/539,151

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L9 ANSWER 41 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1991:82562 CAPLUS
 DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:
                                                            114:82562
114:14124h,14125a
                                                            114:14124h,14125a
Preparation of acyldipeptide amides as tachykinin
antagonists
Matsuo, Masaaki; Hagiwara, Daijiro; Miyake, Hiroshi
Fujisawa Pharmaceutical Co., Ltd., Japan
Eur. Pat. Appl., 13 pp.
CODEN: EEXXDW
 TITLE:
 INVENTOR(S):
PATENT ASSIGNEE(S):
  SOURCE:
 DOCUMENT TYPE:
                                                             English
 LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
           PATENT NO.
                                                             KIND
                                                                            DATE
                                                                                                         APPLICATION NO.
                                                                                                                                                                 DATE
EP 394989 A2 19901031 EP 1990-107822
EP 394989 A3 19910424
EP 394989 B1 19941221
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE
US 5164372 A 19921117 US 1990-505457
CA 2015359 A1 13901028 CA 1990-2015359
JF 03027399 A 19910205 JP 1990-114129
PRIORITY APPLN. INFO.: GB 1989-9795 A
                                                                                                                                                                 19900425
                                                                                                                                                                 19900406
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OTHER SOURCE(S):

AB R1YCOANR2CH(CH2C6H4R3-p)CONR4R5 [R1 = (substituted) alkyl, aryl, arylamino, pyridyl, pyrrolyl, pyrazolopyridyl, quinolyl, Q1, X = CH, N; Z = O, S, NH; R2 = H, alkyl; R3 = H, OH; R4 = (substituted) alkyl; R5 = pyridylalkyl, (substituted) aralkyl; or R4R5 = benzene-condensed alkylene;

A = amino acid residue except D-Trp; Y = bond, alkylene, alkenylene],

prepared Thus, BCC-Q2-Phe-N(Me)CH2Ph [BCC = Me3CO2C, Q2 = (28,4R)-4-hydroxylprolyl residue] (preparation from BCC-Phe-OH given) was deprotected with trifluoroacetic acid and the product was coupled with indole-3-carbonyl chloride (Q3C1) in CH2C12 in the presence of bistrimethylsilylacetamide to give Q3-Q2-Phe-N(Me)CH2Ph. The latter inhibited substance P-induced bronchoconstriction in guinea pigs with an CTC-1-6-0-23 me/ke interstraceally. innibuted substance P-induced bronchoconstriction in guinea pic ED50 of 0.072 mg/kg intratracheally. IT 131948-37-3P RL: BAC (Biological activity or effector, except adverse); BSU (Biological

MARPAT 114:82562

L9 ANSWER 42 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1986:478831 CAPLUS
DOCUMENT NUMBER: 105:78831
OCIGINAL REFERENCE NO: 105:12789a,12792a
3-{2-(Dimethylamino)ethyl]-N-methyl-1H-indole-5-methanesulfonamide
INVENTOR(S): Oxford, Alexander William
Ger. Offen., 57 pp.
CODEN: GWXXBX
DOCUMENT TYPE: Patert

Patent German

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3527648	A1	19860213	DE 1985-3527648	1985080
DE 3527648	C2	19930826		
CH 666026	A.5	19880630	CH 1985-3296	1985073
HU 40077	A2	19861128	HII 1985-2945	1985073
HU 201738	В	19901228		
DK 8503511	A	19860202	DK 1985-3511	1985080
DK 158942	В	19900806		
DK 158942	С	19910121		
FI 8502969	A	19860202	FI 1985-2969	1985080
FI 78466	В	19890428		
FI 78466	C	19890810		
SE 8503680	A	19860202	SE 1985-3680	1985080
SE 452460	В	19871130		
SE 452460	С	19880310		
BE 903006	A1	19860203	BE 1985-215426	1985080
NO 8503046	A	19860203	NO 1985-3046	1985080
NO 164653	В	19900723		
NO 164653	C	19901107		
GB 2162522	A	19860205	GB 1985-19418	1985080
GB 2162522	В	19880224		
AU 8545689	A	19860206	AU 1985-45689	1985080
AU 573878	B2	19880623		
FR 2568571	A1	19860207	FR 1985-11790	1985080
FR 2568571	B1	19880923		
NL 8502171	A	19860303	NL 1985-2171	1985080
NL 188642	В	19920316		
NL 188642	C	19920817		
JP 61047464	A	19860307	JP 1985-168664	1985080
JP 06023197	В	19940330		
ZA 8505818	A	19860430	ZA 1985-5818	1985080
AT 8502266	A	19871215	AT 1985-2266	1985080
AT 386196	В	19880711		
CA 1241004	A1	19880823	CA 1985-487992	1985080
PL 146005	Bl	19881231	PL 1985-254800	1985080
IL 75986	A	19890228	IL 1985-75986	1985080
SU 1498386	AЗ	19890730	SU 1985-3935745	1985080
US 5037845	A	19910806	US 1989-317682	1989030
SK 277952	В6	19950913	SK 1991-4041	1991122
CZ 280530	В6	19960214	CZ 1991-4041	1991122
RITY APPLN. INFO.:			GB 1984-19575	A 1984080
			US 1985-761392	B1 1985080

ANSWER 41 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) study, unclassified); SFN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. of, as tachykinin antagonist) 131948-37-3 CAPLUS 131948-3/-3 CAPLUS
L-Phenylalaninamide,
s-4-hydroxy-1-[(5-hydroxy-1H-indol-3-y1)acetyl]-Lprolyl-N-methyl-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L9 ANSWER 42 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN US 1987-82666 B1 19870807

OTHER SOURCE(S): CASREACT 105:78831

The title compound (I), prepared by 8 methods, is useful in treating

AB The title compound (I), prepared by 8 methods, is useful in treating migraine headaches at 0.1-100 mg per dose, up to 8 times daily. Hydrogenation of 3-(cyanomethyl)-N-methyl-IN-indole-5-methanesulfonamide over prereduced 10% Pd oxide on active C in methanolic and ethanolic MeZNH for 24 h at room temperature gave I (isolated as the succinate). Several formulations were

given. 103628-52-0P

103828-52-0P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and reaction of)
103628-52-0 CAPLUS
H-Indol-8-3-acetamide, N,N-dimethyl-5-[[(methylamino)sulfonyl]methyl](CA INDEX NAME)

L9 ANSWER 43 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1985:560388 CAPLUS DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 103:160388 103:25745a,25748a 103:25745a,25748a
Indole derivatives and their use
Cxford, Alexander William; Evans, Brian; Dowle,
Michael Dennis; Coates, Ian Harold
Glaxo Group Ltd., UK
Ger. Offen., 72 pp.
CODEN: GWXXBX
Patent
German INVENTOR(S): PATENT ASSIGNEE(S): DOCUMENT TYPE: LANGUAGE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 3444572	A1	19850620	DE 1984-3444572		19841206
DE 3444572	C2	19931014	DE 1904-9444972		13041200
FI 8404789	A	19850607	FI 1984-4789		19841205
FI 80260	В	19900131	11 1301 4703		13041600
FI 80260	Č	19900510			
BE 901224	A1	19850606	BE 1984-214125		19841206
DK 8405836	A	19850607	DK 1984-5836		19841206
FR 2555987	A1	19850607	FR 1984-18618		19841206
FR 2555987	B1	19870717	IK 1904-10010		13041200
NO 8404879	A	19850607	NO 1984-4879		19841206
NO 162764	В	19891106	140 1304-4073		13041200
NO 162764	ć	19900214			
SE 8406200	A	19850607	SE 1984-6200		19841206
SE 458446	В	19890403	DE 1304-0200		13041200
SE 458446	ć	19890727			
AU 8436367	A	19850613	AU 1984-36367		19841206
AU 575365	B2	19880728	A0 1904-30307		13041200
NL 8403719	A	19850701	NL 1984-3719		19841206
GB 2150932	A	19850710	GB 1984-30810		19841206
GB 2150932	В	19871028	02 1301 00010		13041100
JP 60155156	A	19850815	JP 1984-258409		19841206
JP 06002733	В	19940112	OF 1904-2304 0 9		17041200
AT 8403873	A	19860515	AT 1984-3873		19841206
AT 381934	В	19861210	2504 0070		15041100
ZA 8409498	A	19860924	ZA 1984-9498		19841206
CH 663411	A5	19871215	CH 1984-5810		19841206
CA 1233183	A1	19880223	CA 1984-469528		19841206
IL 73756	A	19880229	IL 1984-73756		19841206
HU 40624	A2	19870128	ни 1985-2083		19850530
CN 85104233	A	19870107	CN 1985-104233		19850603
CN 85106225	A	19870218	CN 1985-106225		19850819
CN 1015055	В	19911211			
US 4994483	A	19910219	US 1989-443874		19891130
DK 9002140	A	19900906	DK 1990-2140		19900906
JP 03184958	A	19910812	JP 1990-326200		19901129
PRIORITY APPLN, INFO.:			GB 1983-32435	A	19831206
			US 1984-678995	В1	19841206

L9 ANSWER 43 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN US 1987-72786 (Continued) B1 19870713

CASREACT 103:160388; MARPAT 103:160388 OTHER SOURCE(S):

$$\begin{array}{c|c} RR^1NSO_2Z & & H \\ & & \\ &$$

Antimigraine (no data) indolealkanesulfonamides I [R = H, alkyl, alkenyl; R1 = cycloalkyl, Ph, phenylalkyl, R; R2, R3 = H, alkyl, CH2:CHCH2; R2R3 = aralkylidene; Z, Z1 = alkyl-(un)substituted alkylene] were prepared

Thus,

4-02NC6H4CH2CH2SO2Cl was amidated with MeNN2, hydrogenated over Pd-C to the aniline, diazotized, and treated with ZnCl2 to give 4-H2NNHCGH4CH2CH2SO2NHMe. The latter compound was stirred in aqueous MeOH with (MeO)2CH(CH2)3Cl at 50°, NH4OAc added to pH 4, then refluxed 5 h to give 1 (R = Me, Rl-R3 = H, Z = Zl = CH2CH2).

IT 98622-74-3P 98623-48-4P RL EGT (Reactant), SFN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and lithium aluminum hydride reduction of)

RN 98622-74-3 CAPLUS CN 1H-Indole-3-acetamide, N-ethyl-N-methyl-5-[2-[(methylamino)sulfonyl]ethyl]-(CA INDEX NAME)

98623-48-4 CAPLUS 1H-Indole-3-acetamide, N,N-dimethyl-5-[2-[(methylamino)sulfonyl]ethyl]-(CA INDEX NAME)

ANSWER 43 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

L9 ANSWER 44 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 1977:16538 CAPLUS
DOCUMENT NUMBER: 86:16538
ORIGINAL REFERENCE NO.: 86:2689a, 2692a
TITLE: INVENTOR(S): Huebner, Charles F.
PATENT ASSIGNEE(S): Ciba-Geigy A.-G., Switz.
SOURCE: Ger. Offen., 72 pp.
CODEN: GWXXEX
PATENT TYPE: Patent
LANGUAGE: German
TAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2609289	A1	19760930	DE 1976-2609289		19760306
SE 7602729	A	19760913	SE 1976-2729		19760227
NO 7600774	A	19760913	NO 1976-774		19760305
GB 1534351	A	19781206	GB 1976-8902		19760305
FI 7600584	A	19760911	FI 1976-584		19760308
FR 2303541	A1	19761008	FR 1976-6495		19760308
FR 2303541	В1	19791005			
AU 7611750	A	19770915	AU 1976-11750		19760308
IL 49171	A	19781217	IL 1976-49171		19760308
BE 839347	A1	19760909	BE 1976-164977		19760309
DK 7601014	A	19760911	DK 1976-1014		19760309
DK 138893	В	19781113			
DK 138893	C	19790423			
DD 124386	A5	19770216	DD 1976-191763		19760309
NL 7602508	A	19760914	NL 1976-2508		19760310
JP 51113878	A	19761007	JP 1976-26622		19760310
US 4147786	A	19790403	US 1977-797151		19770516
US 4242347	A	19801230	US 1979-50003		19790618
PRIORITY APPLN. INFO.:			US 1975-556600	Α	19750310
			US 1976-654254	АЗ	19760202

OTHER SOURCE(S): CASREACT 86:16538: MARPAT 86:16538

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

AB Indolylethylpiperidines (I; R = e.g., H, 5-C1, 5-Br, 5-F, 7-Me, 7-MeO; R1 = e.g., H, Me; R2 = e.g., H, Me; R3, R4 = e.g., H, H; ethylene, o-phenylene; R5 = e.g., H, Ph; n = 2, 3), useful as antihypertensives, are prepared by various known procedures. Thus, reaction of 3-(2-bromoethyl)indole with 4-ureidopiperidine in DMF 2 days at room temperature in presence of Et3N gives I (R = R1 = R2 = R3 = R4 = R5 = H, n = 2).

IT 61220-26-6P

L9 ANSWER 44 OF 44 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
(prepn. and antihypertensive activity of)
RN 61220-26-6 CAPLUS
CN 2-Imidazolidinone,
1-[1-[2-(6-chioro-iH-indol-3-yl)acetyl]-4-piperidinyl](CA INDEX NAME)